Providing for a State Lottery and administration thereof; authorizing the creation of a State Lottery Commission; prescribing its powers and duties; disposition of funds; violations and penalties therefor; exemption of prizes from State and local taxation and making an appropriation.

Compiler's Note: Section 9 of Act 29 of 1983 provided that Act 91 is repealed insofar as it is inconsistent with Act 29.

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The General Assembly of the Commonwealth of Pennsylvania hereby enacts as follows:
Section 1. Short Title.--(1 deleted by amendment Nov. 21, 1996, P.L.741, No.134)

Section 2. Statement of Purpose.--(2 renumbered 301 and amended Nov. 21, 1996, P.L.741, No.134)

Section 3. Definitions.--(3 deleted by amendment Nov. 21, 1996, P.L.741, No.134)

Section 4. State Lottery Commission; Chairman.--(4 deleted Dec. 15, 1982, P.L.1288, No.291)


Section 6. Powers and Duties of the Secretary of Revenue.--(6 renumbered 303 and amended Nov. 21, 1996, P.L.741, No.134)


Section 7. Lottery Sales Agents; Qualifications; Prohibitions.--(7 renumbered 305 and amended Nov. 21, 1996, P.L.741, No.134)

Section 8. Assignability of Prizes Drawn.--(8 deleted by amendment Nov. 21, 1996, P.L.741, No.134)

Section 9. Sales of Tickets in Excess of Regulated Price; Sales by Non-licensed Persons; Penalties.--(9 renumbered 307 and amended Nov. 21, 1996, P.L.741, No.134)


Section 10. Sales to Certain Persons Prohibited; Penalty.--(10 renumbered 309 and amended Nov. 21, 1996, P.L.741, No.134)

Section 11. Other Laws Inapplicable.--(11 renumbered 310 and amended Nov. 21, 1996, P.L.741, No.134)

Section 12. Disposition of Funds from Sale of Tickets or Shares.--(12 renumbered 311 and amended Nov. 21, 1996, P.L.741, No.134)

Section 12.1. Transportation Assistance Grant.--(12.1 deleted by amendment Nov. 21, 1996, P.L.741, No.134)

Section 13. Exemption of Lottery Prizes from State and Local Taxation.--(13 deleted by amendment, renumbered 312 and amended Nov. 21, 1996, P.L.741, No.134)


Section 16. Appropriation.--(16 deleted by amendment Nov. 21, 1996, P.L.741, No.134)

Section 17. Effective Date.--(17 deleted by amendment Nov. 21, 1996, P.L.741, No.134)

CHAPTER 1
GENERAL PROVISIONS
(Hdg. added Nov. 21, 1996, P.L.741, No.134)

Section 101. Short title.
This act shall be known and may be cited as the State Lottery Law.
(101 added Nov. 21, 1996, P.L.741, No.134)

CHAPTER 3
STATE LOTTERY
(Hdg. added Nov. 21, 1996, P.L.741, No.134)
### Compiler's Note

Section 2 of Act 30 of 1999 provided that as much of Chapter 3 as relates to the payment of administrative expenses, other than the payment of commissions, of Act 91 is repealed.

### Section 301. Statement of purpose.

This chapter is enacted to establish a lottery to be operated by the State, the net proceeds of which are to be used after June 30, 1972, for the purposes of providing property tax relief for the elderly for taxes paid in 1971 and thereafter to persons 65 years of age or older and for providing certain free fixed route local transit services to persons 65 years of age or older and reduced fare on group ride transit service to persons 65 years of age or older. It is further intended to provide a means through which to curb illegal gambling operations in Pennsylvania.

(301 renumbered from 2 and amended Nov. 21, 1996, P.L.741, No.134)

### Section 302. Definitions.

The following words and phrases when used in this chapter shall have the meanings given to them in this section unless the context clearly indicates otherwise:

- "Director." The Director of the Division of the State Lottery.
- "Division." The Division of the State Lottery created by this chapter.
- "Internet instant game." A lottery game in which, by the use of a computer, tablet computer or other mobile device, a player removes the covering from randomly generated numbers or letters which reveal whether the instant ticket is a winning ticket for which money is paid. A player shall not include a retailer who sells or purchases tickets through the lottery central computer.
- "Keno." A lottery game of chance in which a player selects numbers between a preestablished range of numbers with winning numbers selected by a lottery central computer system and displayed on a monitor, commonly known as keno.
- "Lottery" or "State Lottery." The lottery established and operated under this chapter.
- "Secretary." The Secretary of Revenue of the Commonwealth.


### Section 303. Powers and duties of secretary.

(a) Powers and duties enumerated.--In addition to the powers and duties provided by law and the act of April 9, 1929 (P.L.177, No.175), known as The Administrative Code of 1929, the secretary shall have the power and it shall be his duty to operate and administer the lottery, and to promulgate rules and regulations governing the establishment and operation thereof, including, but not limited to:

1. The type of lottery to be conducted. ((1) partially repealed Oct. 30, 2017, P.L.419, No.42)
2. The price, or prices, of tickets or shares in the lottery.
3. The numbers and sizes of the prizes on the winning tickets or shares.
4. The manner of selecting the winning tickets or shares.
5. (i) Except as provided in subparagraph (ii), the manner of payment of prizes to the holders of winning tickets or shares.
   (ii) For new nonmultistate on-line, pari-mutuel games with prizes of $1,000,000 or more and payable in
more than one installment, the player shall have the option at the time of purchase to accept, as full payment of the player's share of the prize won, a lump sum of the prize money allocated to the first place prize category, divided equally by the number of tickets determined by the lottery to be entitled to claim a first place prize, provided that the player specified the lump-sum option at the time of purchase and received a mark on the ticket confirming such request. In cases where the prize pool for the first place prize is not sufficient to fund the guaranteed lump-sum prize as announced by the lottery, the prize pool shall be increased as necessary by the lottery. Players shall be bound by their prewinning choice.

(6) The frequency of the drawings or selections of winning tickets or shares, without limitation.

(7) Without limit as to number, the type or types of locations at which tickets or shares may be sold.

(8) The method to be used in selling tickets or shares, except that sales through the lottery's Internet website or Internet websites operated by licensed lottery retailers of any lottery game or any Internet instant game shall be prohibited unless specifically authorized by law. ((8) amended Oct. 31, 2014, P.L.3041, No.201)

(9) The licensing of agents to sell tickets or shares provided that no person under the age of 21 shall be licensed as an agent.

(10) The manner and amount of compensation, if any, to be paid licensed sales agents necessary to provide for the adequate availability of tickets or shares to prospective buyers and for the convenience of the public.

(11) The apportionment of the total revenues accruing from the sale of lottery tickets or shares and from all other sources among:

(i) the payment of prizes to the holders of winning tickets or shares;

(ii) the payment of costs incurred in the operation and administration of the lottery, including the expenses of the division and the costs resulting from any contract or contracts entered into for promotional, advertising or operational services or for the purchase or lease of lottery equipment and materials; and

(iii) ((iii) deleted by amendment Nov. 21, 1996, P.L.741, No.134)

(iv) for property tax relief and free or reduced fare transit service for the elderly as provided in section 311. For fiscal years beginning before July 1, 2014, no less than 27% of the total revenues accruing from the sale of lottery tickets or shares shall be dedicated to this subparagraph. For fiscal years beginning after June 30, 2014, and ending June 30, 2019, no less than 25% of the total revenues accruing from the sale of lottery tickets or shares shall be dedicated to this subparagraph. For fiscal years beginning after June 30, 2019, and ending June 30, 2024, no less than 20% of the total revenues accruing from the sale of lottery tickets or shares shall be dedicated to this subparagraph. For fiscal years beginning after June 30, 2024, no less than 25% of the total revenues accruing from the sale of lottery tickets or shares shall be dedicated to this subparagraph. ((iv) amended Nov. 27, 2019, P.L.684, No.97)
(11) amended June 30, 2011, P.L.110, No.23)

(11.1) The production and merchandising of promotional items for the lottery.

(12) Such other matters necessary or desirable for the efficient and economical operation and administration of the lottery and for the convenience of the purchasers of tickets or shares and the holders of winning tickets or shares.

(13) The performance of the powers and duties heretofore vested in the State Lottery Commission.

(a.1) Prohibitions.--The secretary may not offer any Internet-based or monitor-based interactive lottery game or simulated casino-style lottery game, including video poker, video roulette, slot machines or video blackjack, through the State Lottery. ((a.1) added Oct. 31, 2014, P.L.3041, No.201)

(b) Reports.--The secretary shall report monthly to the Governor and the Legislature the total lottery revenues, prize disbursements and other expenses for the preceding month, and shall make an annual report, which shall include a full and complete statement of lottery revenues, prize disbursements and other expenses, to the Governor and the Legislature, and including such recommendations for changes in this chapter as the secretary deems necessary or desirable.

(303 renumbered from 6 and amended Nov. 21, 1996, P.L.741, No.134)

Section 304. Commercial advertising.

(a) General rule.--The secretary may enter into contracts with persons, associations or corporations that provide for the placement of commercial advertisements on tickets or shares.

(b) Contracts.--The secretary may enter into the contracts only after completion of the bidding procedure contained in subsection (c).

(c) Bidding procedures.--

(1) The secretary shall, not less than six weeks prior to the date set for opening bids or proposals to place advertisements on the tickets or shares, advertise the opening of proposals for at least three days, the first and last publication to be at least ten days apart, in not fewer than six nor more than 12 newspapers of extensive general circulation in different parts of this Commonwealth. The advertisements shall invite proposals for the placement of commercial advertisements on the tickets or shares, shall direct potential bidders to include with their proposals a specimen advertisement and shall give notice of the time and place where the proposals will be received and when they will be opened.

(2) All proposals shall be delivered to the secretary on or before the hour designated in the invitation to bid, on the day set by the secretary, following the date of the last advertisement, and each bid shall be in duplicates, one of which shall be marked "Duplicate Bid." Each bid shall be enclosed in an envelope, securely sealed, and shall be mailed or delivered to the secretary who shall retain all envelopes unopened until the time fixed for the opening thereof.

(3) The secretary shall, on the date fixed for opening of bids, at the hour designated in the invitation to bid, open and publish the proposals and, as soon thereafter as practicable, award the contract to the highest responsible bidder. The secretary shall have the right to reject any or all bids. The bids, when opened, shall be tabulated and shall be subject to examination by bidders. A record of all bids shall be made by the secretary in a book kept for that purpose.
(4) When no proposal has been received or if for any reason the secretary rejects all proposals, the secretary may advertise again for proposals, giving at least 15 days' notice of the time of receiving the same, which proposals shall be opened, awarded and approved in like manner as the initial bids.

(5) The secretary shall have the discretion to refuse to accept any advertisement that is inappropriate or offensive or displays poor taste. Advertisements for tobacco products or for alcoholic beverages shall not be accepted.

(d) Disposition of revenues.--All revenues derived from contracts entered into under this section shall be deposited in the State Lottery Fund.

(e) Regulations.--The secretary may promulgate rules and regulations to implement the provisions of this section.

(f) Definition.--As used in this section, the term "tickets or shares" shall not include instant game tickets.

(304 renumbered from 6.1 and amended Nov. 21, 1996, P.L.741, No.134)

Section 305. Lottery sales agents.

(a) Licensing.--No license as an agent to sell lottery tickets or shares shall be issued to any person to engage in business exclusively as a lottery sales agent. Before issuing such license the secretary shall consider such factors as:

(1) The financial responsibility and security of the person and his business or activity.
(2) The accessibility of his place of business or activity to the public.
(3) The sufficiency of existing licenses to serve the public convenience.
(4) The volume of expected sales.

(b) Approval of applicant.--If the secretary shall find that the experience, character and general fitness of the applicant are such that the participation of such person as a lottery sales agent will be consistent with the public interest, convenience and necessity, it may thereupon grant a license. Without limiting the generality of the foregoing, the secretary may refuse to issue a license pursuant to this section, or may suspend or revoke a license so issued if it shall find that the applicant or licensee:

(1) Has been convicted of a crime involving moral turpitude.
(2) Has engaged in bookmaking or other form of illegal gambling.
(3) Has been found guilty of any fraud or misrepresentation in any connection.
(4) Has violated any rule, regulation or order of the secretary.

(c) Denial of license.--The secretary may refuse to grant a license or may suspend or revoke a license issued pursuant to this section to a corporation, if it shall determine that:

(1) Any officer, director, member or stockholder of such corporation applying for a license or of any corporation which owns stock in or shares in the profits, or participates in the management of the affairs of such applicant:
(1) has been convicted of a crime involving moral turpitude;
(2) has engaged in bookmaking or other forms of illegal gambling;
(3) has been found guilty of any fraud or misrepresentation in any connection; or
(iv) has violated any rule, regulation or order of the secretary.

(2) The experience, character, or general fitness of any officer, director, or stockholder of any of the aforesaid corporations is such that the participation of such person as a lottery sales agent would be inconsistent with the public interest, convenience or necessity, but if the secretary determines that the interest of any stockholder referred to in this paragraph or in paragraph (1) is sufficient, in the opinion of the secretary, to affect adversely the conduct of a lottery sales agency by such corporation in accordance with the provisions of this chapter, the secretary may disregard such interest in determining whether or not to grant a license to such corporation.

(3) The applicant is not the owner or the lessee of the business at which it will conduct a lottery sales agency pursuant to the license applied for, or that any person, firm, association, or corporation other than the applicant shares, or will share, in the profits of the applicant, other than by dividends as a stockholder, or participates, or will participate, in the management of the affairs of the applicant.

d) Temporary license.—Pending final determination of any question under this section, the secretary may issue a temporary license upon such terms and conditions as it may deem necessary, desirable or proper to effectuate the provisions of this chapter.

e) Resurvey.—Any person who has a pending application for a lottery machine and is currently engaged in the sale of out-of-State lottery tickets may submit a written request to the Department of Revenue for a resurvey. This resurvey shall be completed by the department within 90 days of receipt of the request.

f) Definition.—As used in this section, the term "person" means and includes an individual, association, corporation, club, trust, estate, society, company, joint-stock company, receiver, trustee, assignee, referee or any other person acting in a fiduciary or representative capacity, whether appointed by a court or otherwise, and any combination of individuals. The term shall also mean and include all departments, commissions, agencies and instrumentalities of the State, including counties and municipalities and agencies and instrumentalities thereof.

(305 renumbered from 7 and amended Nov. 21, 1996, P.L.741, No.134)

Section 306. Assignability of prizes drawn.

(a) Assignability.—The right of any person to a prize drawn shall be assignable under the following limited circumstances:

1. Payment of any prize drawn may be paid to the estate of a deceased prize winner.

2. Payment of any prize drawn may be made to any person pursuant to an appropriate judicial order.

3. Payment of any prize drawn may be made to any person pursuant to a voluntary assignment of the right to receive future prize payments, in whole or in part, if the assignment is made to a person or entity designated pursuant to an appropriate judicial order of the court of common pleas located in either the judicial district where the assignor resides or where the division's headquarters are located. Under this paragraph, the court shall issue an order approving the assignment and directing the secretary to pay
the assignee all future prize payments, in whole or in part, if:

(i) the assignment is in writing, executed by the assignor and subject to the laws of this Commonwealth;

(ii) the assignor provides a sworn affidavit to the court attesting that the assignor is of sound mind, is not acting under duress, has been advised regarding the assignment by his or her own independent legal counsel and understands and agrees that, with regard to the assigned payments, the Commonwealth and the secretary shall have no further liability or responsibility to make said payments to the assignor; and

(iii) the proposed assignment does not include or cover payments or portions of payments alleged to be subject to offset under judicial order, unless appropriate provision is made in the order to satisfy the obligations giving rise to the claim for offset, or to offset under any other statute.

(b) Discharge of secretary.--The secretary shall be discharged of all further liability upon payment of a prize pursuant to this section.

(c) Enforcement.--Soliciting or offering rights to lottery prizewinnings, either by assignment or through pledge as collateral for a loan, shall not be deemed selling or offering for sale lottery tickets or shares under this act. Selling or offering for sale assigned or pledged lottery prizewinnings shall not be deemed selling or offering for sale an interest under section 307.

(d) Fees.--The secretary is authorized to establish a reasonable fee to defray any administrative expenses associated with assignments made pursuant to this section, including the cost to the Commonwealth of any processing fee that may be imposed by a private annuity provider. The fee amount shall reflect the direct and indirect costs associated with processing the assignments.

(e) Discharge of Commonwealth.--Upon a voluntary assignment pursuant to appropriate judicial order under subsection (a)(3) of payments due to a prizewinner under a private annuity policy that has been purchased by the lottery for the benefit of a prizewinner, the Commonwealth and the secretary shall be discharged from any and all liability for the payments or portions thereof assigned, and, as to the payments or portions thereof assigned, the assignee shall have recourse only against the private annuity provider and its guarantors and shall have no further recourse against the Commonwealth.

(f) Assignment limitation.--Notwithstanding any other provision of this section, no prizewinner shall have the right to assign prize payments upon:

(1) The issuance by the United States Internal Revenue Service (IRS) of a technical rule letter, revenue ruling or other public ruling of the IRS in which the IRS determines that, based upon the right of assignment provided in this act, a Pennsylvania lottery prizewinner who does not assign any prize payments pursuant to subsection (a)(3) would be subject to an immediate income tax liability for the value of the entire prize rather than annual income tax liability for each installment when paid.

(2) The issuance by a court of competent jurisdiction of a published decision holding that, based upon the right of assignment provided in this act, a Pennsylvania lottery prizewinner who does not assign any prize payments pursuant to subsection (a)(3) would be subject to an immediate income
tax liability for the value of the entire prize rather than annual income tax liability for each installment when paid.

(g) Filing of letter decision.--Upon receipt of a letter or ruling from the IRS or a published decision of a court of competent jurisdiction, as specified in subsection (f), the executive director shall immediately file a copy of that letter, ruling or published decision with the Secretary of State. Immediately upon the filing by the director of a letter, ruling or published decision with the Secretary of State, a prizewinner shall be ineligible to assign a prize pursuant to subsection (a)(3).

(306 added Nov. 21, 1996, P.L.741, No.134)

Section 307. Ticket sales.
(a) Prices.--No person shall sell, resell or engage in the business of reselling lottery tickets or shares at a price greater than that fixed by rule or regulation of the department. Price shall include any fee associated with the acquisition or transportation of lottery tickets or shares.
(b) Construction.--No person other than a licensed lottery sales agent shall sell lottery tickets or shares, except that nothing in this section shall be construed to prevent any person from giving lottery tickets or shares to another as a gift.
(c) Lotteries of other states.--Except as provided in this chapter, no person shall engage in the sale or offering for sale within this Commonwealth of any interest in a lottery of another state or government whether or not such interest is an actual lottery ticket, receipt, contingent promise to pay, order to purchase or other record of such interest.
(d) Penalty.--Any person convicted of violating this section shall be guilty of a misdemeanor and upon conviction thereof, shall be sentenced to pay a fine of not more than $2,000.

(307 renumbered from 9 and amended Nov. 21, 1996, P.L.741, No.134)

Section 308. Compact to sell tickets.
The secretary shall enter into a compact with any other states that permit sale of Pennsylvania lottery tickets within their borders to sell those states' lottery tickets within this Commonwealth.

(308 renumbered from 9.1 and amended Nov. 21, 1996, P.L.741, No.134)

Section 309. Certain sales prohibited.
(a) Minors.--
(1) No ticket or share shall be sold to any person under 18 years of age. For the purpose of making a gift, a person 18 years of age or older may purchase a ticket or share for the benefit of a person less than that age.
(2) Any agent or employee of any agent who knowingly sells a lottery ticket or share to any person under 18 years of age shall be guilty of a misdemeanor, and upon conviction thereof, shall be sentenced to pay a fine of not more than $500.
(b) Certain employees.--No ticket or share shall be sold to and no prize shall be awarded to any officer or employee of the division in the Department of Revenue or any spouse, child, brother, sister or parent residing as a member of the same household in the principal place of abode of any of the foregoing persons.

(309 renumbered from 10 and amended Nov. 21, 1996, P.L.741, No.134)

Section 310. Other laws inapplicable.
No other law providing any penalty or disability for the sale of lottery tickets or shares or any acts done in connection
with a lottery shall apply to the sale of tickets or shares or acts performed pursuant to this chapter.  

(310 renumbered from 11 and amended Nov. 21, 1996, P.L.741, No.134)

Section 311. Disposition of funds.  

(a) State Lottery Fund.--All moneys received from the operation of the State lottery shall be deposited in a State Lottery Fund which is hereby created. Such moneys shall be used to the extent necessary for the payment of lottery prizes but the amount so used shall not be less than 40% of the amount of which tickets or shares have been sold. All payments of lottery prizes and for expenses of operation of the lottery shall be made as provided by law. All moneys remaining after payment of prizes and operating expenses shall remain in the State Lottery Fund and shall be allocated for the purpose of providing property tax relief for the elderly for taxes paid in 1971 and thereafter pursuant to the provisions of the act of March 11, 1971 (P.L.104, No.3), known as the Senior Citizens Rebate and Assistance Act, and for the purpose of providing free or reduced fare transit service for the elderly pursuant to Chapter 9 and the act of February 11, 1976 (P.L.14, No.10), known as the Pennsylvania Rural and Intercity Common Carrier Surface Transportation Assistance Act. In the event sufficient funds are not available from the lottery receipts to meet the requirements of the Senior Citizens Rebate and Assistance Act or for providing free or reduced fare transit service for the elderly under Chapter 9 and the Pennsylvania Rural and Intercity Common Carrier Surface Transportation Assistance Act, additional funds to fulfill these obligations shall be appropriated from the General Fund for this purpose.  

(b) Appropriations.--The moneys in said State Lottery Fund shall be appropriated only:  

(1) For the payment of prizes to the holders of winning lottery tickets or shares.  

(2) For the expenses of the division in its operation of the lottery.  

(3) For property tax relief and free or reduced fare transit service for the elderly as provided under subsection (a).  

(311 renumbered from 12 and amended Nov. 21, 1996, P.L.741, No.134)

Section 312. Tax exemption.  

No State or local taxes of any kind whatsoever shall be imposed upon the proceeds from a prize awarded by the State lottery.  

(312 renumbered from 13 and amended Nov. 21, 1996, P.L.741, No.134)

Compiler's Note: Section 50(2) of Act 84 of 2016 provided that section 312 is repealed insofar as it is inconsistent with Act 84.

Section 313. Unclaimed prize money.  

Unclaimed prize money on a winning lottery ticket or share shall be retained by the secretary for payment to the person entitled thereto for one year after the drawing in which the prize was won. If no claim is made within such period, the prize money shall be paid into the State Lottery Fund and used for purposes as otherwise herein provided.  

(313 renumbered from 14 and amended Nov. 21, 1996, P.L.741, No.134)

Section 314. Deposits and transactions.
The secretary may, in his discretion, require any or all lottery sales agents to deposit to the credit of the State Lottery Fund in banks, designated by the State Treasurer, all moneys received by such agents from the sale of lottery tickets or shares, less the amount, if any, retained as compensation for the sale of the tickets or shares, and to file with the secretary or his designated agents reports of their receipts and transactions in the sale of lottery tickets in such form and containing such information as he may require. The secretary may make such arrangements for any person, including a bank, to perform such functions, activities or services in connection with the operation of the lottery as he may deem advisable pursuant to this chapter and the rules and regulations of the department, and such functions, activities or services shall constitute lawful functions, activities and services of such person.

(314 renumbered from 15 and amended Nov. 21, 1996, P.L.741, No.134)

Section 315. Report.

The Department of Revenue shall submit a report to the Governor, the chairman and minority chairman of the Appropriations Committee of the Senate, the chairman and minority chairman of the Appropriations Committee of the House of Representatives, the chairman and minority chairman of the Aging and Youth Committee of the Senate, the chairman and minority chairman of the Aging and Older Adult Services Committee of the House of Representatives, the chairman and minority chairman of the Finance Committee of the Senate and the chairman and minority chairman of the Finance Committee of the House of Representatives by September 1 of each year. The report shall set forth current lottery profits and the State Lottery's plan for increasing future profits. This report shall be posted on the department's publicly accessible Internet website.


SUBCHAPTER A
PRELIMINARY PROVISIONS
(Hdg. added July 7, 2006, P.L.1061, No.111)

Section 501. Legislative findings.

Finding that an increasing number of the Commonwealth's elderly citizens who are living on fixed incomes are experiencing difficulties in meeting the costs of life-sustaining prescription drugs, the General Assembly, in its responsibilities to provide for the health, welfare and safety of the residents of this Commonwealth, hereby continues a limited State pharmaceutical assistance program for the elderly.

(501 added Nov. 21, 1996, P.L.741, No. 134)

Section 502. Definitions.

The following words and phrases when used in this chapter shall have the meanings given to them in this section unless the context clearly indicates otherwise:

"A-rated generic therapeutically equivalent drug." A drug product that the Commissioner of Food and Drugs of the United States Food and Drug Administration has approved as safe and effective and has determined to be therapeutically equivalent, as listed in "The Approved Drug Products with Therapeutic Equivalence Evaluations" (Food and Drug Administration "Orange Book"), with a specific "A" code designation only.
"Average wholesale cost." The cost of a dispensed drug based upon the price published in a national drug pricing system in current use by the Department of Aging as the average wholesale price of a prescription drug in the most common package size.

"Average wholesale price." Average wholesale cost.

"Board." The Pharmaceutical Assistance Advisory Board. (Def. amended Oct. 23, 2018, P.L.579, No.87)

"CMS." The Centers for Medicare and Medicaid Services of the United States. (Def. added Nov. 26, 2003, P.L.212, No.37)

"Claimant." An eligible person who is enrolled in the program. (Def. added July 7, 2006, P.L.1061, No.111)

"DESI." The Drug Efficacy Study Implementation List. (Def. added Nov. 26, 2003, P.L.212, No.37)

"Department." The Department of Aging of the Commonwealth.

"Eligible person." A resident of the Commonwealth for no less than 90 days, who is 65 years of age or older, whose annual income is less than the maximum annual income and who is not otherwise qualified for public assistance under the act of June 13, 1967 (P.L.31, No.21), known as the Public Welfare Code. (Def. amended July 7, 2006, P.L.1061, No.111)

"FDA." The United States Food and Drug Administration of the Public Health Service of the Department of Health and Human Services.

"HCFA." (Def. deleted by amendment Nov. 26, 2003, P.L.212, No.37)

"Health maintenance organization." An organized system which combines the delivery and financing of health care and which provides basic health services to voluntarily enrolled subscribers for a fixed prepaid fee. (Def. added Nov. 26, 2003, P.L.212, No.37)

"Income." All income from whatever source derived, including, but not limited to, salaries, wages, bonuses, commissions, income from self-employment, alimony, support money, cash public assistance and relief, the gross amount of any pensions or annuities, including railroad retirement benefits, all benefits received under the Social Security Act (49 Stat. 620, 42 U.S.C. § 301 et. seq.) net of amounts withheld for Medicare Part B premium payment, all benefits received under State unemployment insurance laws, all interest received from the Federal Government or any state government or any instrumentality or political subdivision thereof, realized capital gains, rentals, workmen's compensation and the gross amount of loss of time insurance benefits, life insurance benefits and proceeds, except the first $10,000 of the total of death benefits payments, and gifts of cash or property, other than transfers by gift between members of a household, in excess of a total value of $300, but shall not include surplus food or other relief in kind supplied by a government agency or property tax rebate nor shall the term include any State veterans' benefit payments. (Def. amended Oct. 29, 2020, P.L.717, No.82)

"LEP" or "late enrollment penalty." The amount added to the Part D plan premium of either:

(1) an individual who did not obtain creditable prescription drug coverage as defined under 42 CFR § 423.56 (relating to procedures to determine and document creditable status of prescription drug coverage) when the individual was first eligible for Part D; or

(2) an individual who had a break in creditable prescription drug coverage of at least 63 consecutive days. The LEP is considered a part of the plan premium. (Def. added Oct. 23, 2018, P.L.579, No.87)
"Less expensive." The lowest net cost to the program. The net cost shall include the amount paid by the Commonwealth to a pharmacy for a drug under a current retail pharmacy reimbursement formula less any discount or rebates, including those paid during the previous calendar quarter and inclusive of all dispensing fees. (Def. added Nov. 30, 2004, P.L.1722, No.219)

"Maintenance medication." A medication prescribed for a chronic, long-term condition and taken on a regular, recurring basis. (Def. added Oct. 23, 2018, P.L.579, No.87)

"Maximum annual income." For PACE eligibility, the term shall mean annual income which shall not exceed $14,500 in the case of single persons nor $17,700 in the case of the combined annual income of persons married to each other. For PACENET eligibility, the term shall mean the annual income limits established under section 519. Persons may, in reporting income to the Department of Aging, round the amount of each source of income and the income total to the nearest whole dollar, whereby any amount which is less than 50¢ is eliminated. (Def. amended July 7, 2006, P.L.1061, No.111)

"Medicare advantage." A plan of health benefits coverage offered under a policy, contract or plan by an organization certified under 42 U.S.C. § 1395w-26 (relating to establishment of standards) and formerly referred to as Medicare+Choice. (Def. added July 7, 2006, P.L.1061, No.111)


"Medication synchronization." The coordination of prescription drug filling or refilling by a pharmacy or dispensing physician for a program participant taking two or more medications for the purpose of improving medication adherence. (Def. added Oct. 23, 2018, P.L.579, No.87)

"PACE." The Pharmaceutical Assistance Contract for the Elderly program provided for in this chapter.

"PACENET." The Pharmaceutical Assistance Contract for the Elderly Needs Enhancement Tier provided for in this chapter.


"Part D plan" or "PDP." A prescription drug plan approved under the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (Public Law 108-173, 117 Stat. 2066) in the PDP region that includes this Commonwealth and approved by the Department of Aging of the Commonwealth and the Centers for Medicare and Medicaid Services of the United States for coordination of benefits with the programs established under this chapter. (Def. added July 7, 2006, P.L.1061, No.111)

"Pharmacy." A pharmacy licensed by the Commonwealth.

"Preferred provider organization." An entity organized and operating under 40 Pa.C.S. Ch. 63 (relating to professional health services plan corporations). (Def. added Nov. 26, 2003, P.L.212, No.37)

"Prescription drug." All drugs requiring a prescription in this Commonwealth, insulin, insulin syringes and insulin needles. Experimental drugs or drugs prescribed for wrinkle removal or hair growth are prohibited.
"Private contractor." A person, partnership or corporate entity who enters into a contract with the Commonwealth to provide services under the provisions of this chapter.

"Program." The Pharmaceutical Assistance Contract for the Elderly (PACE) and the Pharmaceutical Assistance Contract for the Elderly Needs Enhancement Tier (PACENET) as established by this chapter. (Def. amended July 7, 2006, P.L.1061, No.111)

"Provider." A pharmacy, dispensing physician or certified registered nurse practitioner enrolled as a provider in the program. (Def. amended Nov. 26, 2003, P.L.212, No.37)

"Regional benchmark premium." The average Part D premium calculated annually by the Centers for Medicare and Medicaid Services of the United States for PDPs in the PDP region that includes this Commonwealth. (Def. added July 7, 2006, P.L.1061, No.111)

"State veterans' benefit payments." Service-connected compensation or payments provided to a veteran or an unmarried surviving spouse of a veteran by a State agency or authorized under State law. (Def. added Oct. 29, 2020, P.L.717, No.82)

(502 added Nov. 21, 1996, P.L.741, No.134)

Compiler's Note: The short title of the act of June 13, 1967 (P.L.31, No.21), known as the Public Welfare Code, referred to in this section, was amended by the act of December 28, 2015 (P.L.500, No.92). The amended short title is now the Human Services Code.

Compiler's Note: See section 14 of Act 111 of 2006 in the appendix to this act for special provisions relating to eligibility for PACE or PACENET program.

SUBCHAPTER B
PROGRAMS
(Hdg. added July 7, 2006, P.L.1045, No.111)

Section 503. Determination of eligibility.
(a) Duties of department.--The department shall adopt regulations relating to the determination of eligibility of prospective claimants and providers, including dispensing physicians and certified registered nurse practitioners when acting in accordance with rules and regulations promulgated by the State Board of Nursing as required by the act of May 22, 1951 (P.L.317, No.69), known as The Professional Nursing Law, and the State Board of Pharmacy minimum standards of practice, and the determination and elimination of program abuse. To this end, the department shall establish a compliance unit staffed sufficiently to fulfill this responsibility. The department shall have the power to declare ineligible any claimant or provider who abuses or misuses the established prescription plan. The department shall have the power to investigate cases of suspected provider or recipient fraud.
(b) Social Security cost-of-living adjustment.--
(1) Notwithstanding any other provision of this act to the contrary, persons who, as of December 31, 2020, are enrolled in the PACENET program shall remain eligible for the PACENET program if the maximum income limit is exceeded due solely to a Social Security cost-of-living adjustment.
(2) Notwithstanding any other provision of this act to the contrary, persons who, as of December 31, 2020, are enrolled in the PACE program shall remain eligible for the PACE program if the maximum income limit is exceeded due solely to a Social Security cost-of-living adjustment.
Eligibility in the PACE or PACENET program pursuant to this subsection shall expire on December 31, 2023.
(503 amended July 9, 2008, P.L.934, No.69)

Compiler's Note: Section 1.1 of Act 21 of 2011, which reenacted and amended subsec. (b), provided that the reenactment and amendment of subsec. (b) shall apply retroactively to December 30, 2010.
Section 14 of Act 37 of 2003, which amended section 503, provided that the Department of Aging may use a PACE or PACENET program applicant's most recent annual income information to determine program eligibility until April 1, 2004.

Section 504. Physician, certified registered nurse practitioner and pharmacy participation.

Any physician, certified registered nurse practitioner, pharmacist, pharmacy or corporation owned in whole or in part by a physician, certified registered nurse practitioner or pharmacist enrolled as a provider in the program or who has prescribed medication for a claimant who is precluded or excluded for cause from the Department of Public Welfare's Medical Assistance Program shall be precluded or excluded from participation in the program. No physician or certified registered nurse practitioner precluded or excluded from the Department of Public Welfare's Medical Assistance Program shall have claims resulting from prescriptions paid for by the program.
(504 amended July 7, 2006, P.L.1061, No.111)

Compiler's Note: The Department of Public Welfare, referred to in this section, was redesignated as the Department of Human Services by Act 132 of 2014.

Section 505. Drug utilization review system.

(a) Establishment.--The department shall ensure that a state-of-the-art therapeutic drug utilization review system is established to monitor and correct misutilization of drug therapies.

(b) Review.--The department shall review utilization data provided from a PDP to monitor increases in drug utilization among claimants and determine if disease management intervention is needed.
(505 amended July 7, 2006, P.L.1061, No.111)

Section 506. Reduced assistance.

Any claimant whose prescription drug costs are covered in part by any other plan of assistance or insurance, including Part D, may be required to receive reduced assistance under the provisions of this subchapter or be subject to coordination of benefits under this chapter.
(506 amended July 7, 2006, P.L.1045, No.111)

Section 507. Rebates for expenses prohibited.

A system of rebates or reimbursements to the claimant for prescription drugs shall be prohibited.
(507 added Nov. 21, 1996, P.L.741, No.134)

Section 508. Request for proposal.

(a) General rule.--The department shall prepare a request for proposal for the purpose of providing pharmaceutical assistance for the elderly within this Commonwealth. Upon the adoption of the General Fund budget, the Department of Revenue shall be authorized to transmit the appropriated funds in the State Lottery Fund to the State Treasurer to be deposited in the Pharmaceutical Assistance Contract for the Elderly Fund.
This fund shall consist of appropriations and interest and shall be created by the State Treasurer to fund the operations of the program by the department and the private contractor. Funds not expended in the fiscal year in which they were appropriated shall not lapse and be available for use in the next fiscal year.

(b) Additional requests for proposals.--To provide for the continued operation of the program, the department shall prepare, as needed, requests for proposals, in addition to that set forth in subsection (a), for the purpose of providing pharmaceutical assistance for the elderly within this Commonwealth. A request for proposal shall require potential private contractors to submit a proposal for a period of time and with monetary limitations as determined by the department. Upon the enactment of an appropriation from the State Lottery Fund, the Department of Revenue shall be authorized to transmit the appropriated amount to the State Treasurer to be deposited in the Pharmaceutical Assistance Contract for the Elderly Fund. Funds not expended in the fiscal year in which they were appropriated shall not lapse and shall be available for use in the next fiscal year.

(508 added Nov. 21, 1996, P.L.741, No.134)

Section 509. Program generally.
The program shall include the following:
(1) Participating pharmacies shall be paid within 21 days of the contracting firm receiving the appropriate substantiation of the transaction. Pharmacies shall be entitled to interest for payment not made within the 21-day period at a rate approved by the board.
(2) Collection of the copayment by pharmacies shall be mandatory.
(3) Claimants are not required to maintain records of each transaction.
(4) A system of rebates or reimbursements to claimants for pharmaceutical expenses shall be prohibited.
(5) PACE shall include participant copayment schedules for each prescription, including a copayment for generic or multiple-source drugs that is less than the copayment for single-source drugs. The department shall annually calculate the copayment schedules based on the Prescription Drugs and Medical Supplies Consumer Price Index. When the aggregate impact of the Prescription Drugs and Medical Supplies Consumer Price Index equals or exceeds $1, the department shall adjust the copayment schedules. Each copayment schedule shall not be increased by more than $1 in a calendar year.
(6) ((6) repealed Nov. 21, 2016, P.L.1318, No.169)
(7) In no case shall the Commonwealth or any claimant be charged more than the price of the drug at the particular pharmacy on the date of the sale.
(8) The Governor may, based upon certified State Lottery Fund revenue that is provided to both the chairman and minority chairman of the Appropriations Committee of the Senate and the chairman and minority chairman of the Appropriations Committee of the House of Representatives, and after consultation with the board, decrease the eligibility limits established in this subchapter.

(509 amended July 7, 2006, P.L.1061, No.111)

Section 510. Generic drugs.
(a) In general.--((a) repealed Nov. 21, 2016, P.L.1318, No.169)
(b) Generic not accepted.--((b) repealed Nov. 21, 2016, P.L.1318, No.169)
(c) Generic drugs not deemed incorrect substitution.--The dispensing of an A-rated generic therapeutically equivalent drug in accordance with this subchapter shall not be deemed incorrect substitution under section 6(a) of the Generic Equivalent Drug Law.

(d) Medical exception.--A medical exception process shall be established by the department, which shall be published as a notice in the Pennsylvania Bulletin and distributed to providers and recipients in the program.

(510 amended July 7, 2006, P.L.1061, No.111)

Section 511. Supply.

Prescription benefits for any single prescription shall be limited to a 30-day supply of the prescription drug or 100 units, whichever is less, except that, in the case of diagnosis for acute conditions, the limitation shall be a 15-day supply. This limitation shall not apply to topical ointments or gels that are not available in containers which meet the size and supply restrictions set forth in this section.

(511 added Nov. 21, 1996, P.L.741, No.134)

Section 512. Formulary.

The department may establish a formulary of the drugs which will not be reimbursed by the program. This formulary shall include experimental drugs and drugs on the Drug Efficacy Study Implementation List prepared by CMS. A medical exception may be permitted by the department for reimbursement of a drug on the Drug Efficacy Study Implementation List upon declaration of its necessity on the prescription by the treating physician or certified registered nurse practitioner, except that, for DESI drugs for which the FDA has issued a Notice for Opportunity Hearing (NOOH) for the purpose of withdrawing the New Drug Application approved for that drug, reimbursement coverage shall be discontinued under the provisions of this subchapter.

(512 amended July 7, 2006, P.L.1061, No.111)

Section 513. Mail order system.

(a) The department may not enter into a contract with a private contractor for an exclusive mail-order system for the delivery of prescription drugs under this program. Only mail-order pharmacy services provided by pharmacies which are licensed by the Commonwealth and which have their principal place of business within this Commonwealth may participate as providers under the program. The department shall develop and promulgate specific regulations governing the practice of mail-order pharmacy and other enrolled providers to include the following minimum standards of practice to ensure the health, safety and welfare of program participants:

(1) The appropriate method or methods by which such pharmacies shall verify the identity of the program recipient and the authenticity of prescriptions received.

(2) The appropriate method or methods by which such pharmacies shall mail or deliver prescription drugs to program recipients ensuring, to the maximum extent possible, that the intended program recipient is the actual ultimate recipient of any prescription dispensed by such pharmacies.

(3) The appropriate method or methods by which such pharmacies shall communicate with program participants in emergency situations.

(b) Notwithstanding any provision of law to the contrary, a claimant may use any and all pharmacy services offered by a PDP or Medicare Advantage Prescription Drug Plan to receive drugs and shall be permitted to continue to use those services throughout the noncoverage phase.
(c) Nothing in this section shall require a claimant to use mail-order services.

Section 514. Indication of price.
The retail price of the prescription shall be indicated on the label of the prescription container or furnished by separate receipt.

Section 515. Reimbursement.
For-profit third-party insurers, health maintenance organizations, preferred provider organizations, not-for-profit prescription plans, Medicare advantage plans and PDPs shall be responsible for any payments made to a providing pharmacy on behalf of a claimant covered by such a third party. Final determination as to the existence of third-party coverage shall be the responsibility of the department.

Section 515.1. Medication synchronization.
(a) Prorated daily cost-sharing fees.--The program shall permit and apply a prorated daily cost-sharing fee to prescription drugs that are dispensed by a pharmacy for less than a 30 days' supply if the pharmacist or prescriber determines the fill or refill to be in the best interest of the program participant and the program participant requests or agrees to less than a 30 days' supply for the purpose of medication synchronization. The program may not use payment structures incorporating prorated dispensing fees.

(b) Full payment.--Dispensing fees for a partial supply or refilled prescription shall be paid in full for each maintenance medication dispensed, regardless of any prorated copay for the beneficiary or fee paid for alignment services.

(c) Partial supply.--The program may not deny coverage for the dispensing of a maintenance medication that is dispensed by a network pharmacy on the basis that the dispensing is for a partial supply if the prescriber or pharmacist determines the fill or refill is in the best interest of the patient and the patient requests or agrees to a partial supply for the purpose of medication synchronization.

(d) Annual limitation.--The fill or refill under this section shall be limited to three times a year for each maintenance medication for a covered individual. For each clinically necessary synchronization thereafter, approval may be required at the discretion of the program.

(e) Override of denial codes.--Subject to section 520(c.1), the program shall allow a pharmacy to override any denial codes indicating that a prescription drug is being refilled too soon for the purposes of medication synchronization utilizing the submission clarification and message codes as adopted by the National Council for Prescription Drug Programs or alternative codes provided by the program.

(f) Exemption.--This section does not apply to prescription drugs that are either:
(1) unit-of-use packaging for which medication synchronization is not possible; or
(2) controlled substances classified in Schedule II under section 4(2) of the act of April 14, 1972 (P.L.233, No.64), known as The Controlled Substance, Drug, Device and Cosmetic Act.

Section 516. Nonliability.
(a) General rule.--Any person rendering service as a member of a utilization review committee for this program shall not
be liable for any civil damages as a result of any acts or omissions in rendering the service as a member of any such committee except any acts or omissions intentionally designed to harm or any grossly negligent acts or omissions which result in harm to the person receiving such service.

(b) Department personnel.--Any officer or employee of the department rendering service as a member of a utilization review committee for this program shall not be liable for any civil damages as a result of any acts or omissions in rendering the service as a member of any such committee or as a result of any decision or action in connection with the program except any acts or omissions intentionally designed to harm or any grossly negligent acts or omissions which result in harm to the person receiving such service.

(516 amended Nov. 26, 2003, P.L.212, No.37)

Section 517. Income verification.

(a) Procedure.--The department shall annually verify the income of claimants. The department shall verify the income of claimants by requiring income documentation from the claimants. An application for benefits under this subchapter shall constitute a waiver to the department of all relevant confidentiality requirements relating to the claimant's Pennsylvania State income tax information in the possession of the Department of Revenue. The Department of Revenue shall provide the department with the necessary income information shown on the claimant's Pennsylvania State income tax return solely for income verification purposes.

(b) Information confidential.--It shall be unlawful for any officer, agent or employee of the department to divulge or make known in any manner whatsoever any information gained through access to the Department of Revenue information except for official income verification purposes under this subchapter or as authorized under section 535.

(c) Penalty.--A person who violates this section commits a misdemeanor and shall, upon conviction, be sentenced to pay a fine of not more than $1,000 or to imprisonment for not more than one year, or both, together with the cost of prosecution, and, if the offender is an officer or employee of the Commonwealth, he shall be dismissed from office or discharged from employment.

(d) Coordination with Department of Public Welfare.--To the extent possible, the department and the Department of Public Welfare shall coordinate efforts to facilitate the application and enrollment of eligible older people in the Medicaid Healthy Horizons Program by processing these applications at senior citizens centers and other appropriate facilities providing services to the elderly.

(e) Coordination with the Department of Health.--The department shall establish a method to be used at least once each calendar month, to cross-reference the department's roster of claimants with the death records information from the Department of Health. A claimant who is found to have a death record shall be subject to an immediate cancellation of benefits. ((e) added April 17, 2020, P.L.81, No.14)

(517 amended July 7, 2006, P.L.1061, No.111)

Compiler's Note: The Department of Public Welfare, referred to in this section, was redesignated as the Department of Human Services by Act 132 of 2014.
subchapter. The department shall select a proposal that
includes, but is not limited to, the criteria set forth in this
subchapter.

(518 amended July 7, 2006, P.L.1061, No.111)

Section 519. The Pharmaceutical Assistance Contract for the
Elderly Needs Enhancement Tier.

(a) Establishment. -- There is hereby established within the
department a program to be known as the Pharmaceutical
Assistance Contract for the Elderly Needs Enhancement Tier
(PACENET).

(b) PACENET eligibility. -- A person with an annual income
of not less than $14,500 and not more than $33,500 in the case
of a single person and of not less than $17,700 and not more
than $41,500 in the case of the combined income of persons
married to each other shall be eligible for enhanced
pharmaceutical assistance under this section. A person may, in
reporting income to the department, round the amount of each
source of income and the income total to the nearest whole
dollar, whereby any amount which is less than 50¢ is eliminated.
((b) amended Dec. 22, 2021, P.L.454, No.94)

(c) Deductible. -- ((c) deleted by amendment July 7, 2006,
P.L.1045, No.111)

(c.1) Premium. -- In those instances in which a PACENET
claimant is not enrolled in Part D pursuant to section 533, the
claimant shall be required to pay a monthly premium equivalent
to the regional benchmark premium for each month the claimant
is dispensed a prescription drug. The claimant shall not be
required to pay a monthly premium for any month the claimant
is not dispensed a prescription drug. ((c.1) amended Dec. 22,
2021, P.L.454, No.94)

(d) Copayment. --

(1) For claimants under this section, the copayment
schedule shall be:

(i) eight dollars for noninnovator multiple source
drugs as defined in section 702; or

(ii) fifteen dollars for single-source drugs and
innovator multiple-source drugs as defined in section
702.

(2) The department shall annually calculate the
copayment schedules based on the Prescription Drugs and
Medical Supplies Consumer Price Index. When the aggregate
impact of the Prescription Drugs and Medical Supplies
Consumer Price Index equals or exceeds $1, the department
shall adjust the copayment schedules. Each copayment schedule
shall not be increased by more than $1 in a calendar year.
(519 amended July 7, 2006, P.L.1061, No.111)

Compiler's Note: Section 4 of Act 219 of 2004 provided that,
notwithstanding any other provision of law to the
contrary, persons, who, as of December 31, 2004, are
enrolled in the PACENET program established pursuant to
section 519 shall remain eligible for the PACENET program
if the maximum income limit is exceeded due solely to a
Social Security cost-of-living adjustment. Eligibility
in the PACENET program pursuant to section 44 shall
expire on December 31, 2005.

Compiler's Note: Section 14 of Act 37 of 2003, which amended
section 519, provided that the Department of Aging may
use a PACE or PACENET program applicant's most recent
annual income information to determine program
eligibility until April 1, 2004.

Section 520. Board.
(a) Establishment.--The Pharmaceutical Assistance Advisory Board is continued to ensure that the program is providing and continues to provide the assistance intended in a fiscally responsible manner without excessively hampering the pharmaceutical industry.

(b) Composition.--The board shall be comprised of the following persons:

1. The Secretary of Aging, who shall serve as its chairman.
2. The Secretary of Revenue.
3. The Secretary of Health.
4. Nine public members appointed as follows:
   i. Four practicing Pennsylvania pharmacists whose names are jointly submitted by the Pennsylvania Pharmacists Association and the Pennsylvania Association of Chain Drug Stores and then appointed by the following:
      A. One member appointed by the President pro tempore of the Senate.
      B. One member appointed by the Minority Leader of the Senate.
      C. One member appointed by the Speaker of the House of Representatives.
      D. One member appointed by the Minority Leader of the House of Representatives.
   ii. Five individuals appointed by the Governor that include the following:
      A. One representative from the pharmaceutical industry.
      B. Four senior citizens who have not been a part of the pharmaceutical industry, two of whom may be senior advocates.
5. Should a board vacancy not be filled by the appointing authority within 60 days, the power to appoint an individual to the vacancy shall be given to the Secretary of Aging.

(c) Review.--Using the annual report submitted by the department pursuant to section 2102 and other appropriate data sources, the board shall conduct an annual review. The board shall develop recommendations concerning any changes in the level of copayment or deductible. The board shall review the department's therapeutic drug utilization review program on an ongoing basis. The board may also recommend other changes in the structure of the program and direct the department to enter into discussions with the private contractor concerning amendments to the contract, or the department may enter into such discussions at its discretion. The copayment or deductible schedule shall only be adjusted on an annual basis.

(c.1) Powers and duties.--The board shall advise on the following:

1. The development and implementation of medication synchronization and medication therapy management programs and reimbursement methodologies.
2. Adjustment of the dispensing fee, as needed.
3. Cost-of-living adjustment increases for medication synchronization, medication therapy management and the dispensing fee.
4. The development and modernization of the program, as necessary, to ensure that the program is providing and continues to provide the assistance intended in a fiscally responsible manner. (4) added Dec. 22, 2021, P.L.454, No.94)

(d) Meetings.--The board shall meet at least two times per year and not more than four times per year.
(520 amended Oct. 23, 2018, P.L.579, No.87)
Section 520.1. Pharmacy best practices and cost controls review. (520.1 deleted by amendment July 7, 2006, P.L.1061, No.111)
Section 521. Penalties.
(a) Prohibited acts.--It shall be unlawful for any person to submit a false or fraudulent claim or application under this subchapter, including, but not limited to:
   (1) aiding or abetting another in the submission of a false or fraudulent claim or application;
   (2) receiving benefits or reimbursement under a private, Federal or State program for prescription assistance and claiming or receiving duplicative benefits hereunder;
   (3) soliciting, receiving, offering or paying any kickback, bribe or rebate, in cash or in kind, from or to any person in connection with the furnishing of services under this subchapter;
   (4) engaging in a pattern of submitting claims that repeatedly uses incorrect National Drug Code numbers; or
   (5) otherwise violating any provision of this subchapter.
(b) Civil penalty.--In addition to any appropriate criminal penalty for prohibited acts under this subchapter whether or not that act constitutes a crime under 18 Pa.C.S. (relating to crimes and offenses), a provider who violates this section may be liable for a civil penalty in an amount not less than $500 and not more than $10,000 for each violation of this act which shall be collected by the department. Each violation constitutes a separate offense. If the department collects three or more civil penalties against the same provider, the provider shall be ineligible to participate in either PACE or PACENET for a period of one year. If more than three civil penalties are collected from any provider, the department may determine that the provider is permanently ineligible to participate in PACE or PACENET.
(c) Suspension of license.--The license of any provider who has been found guilty under this subchapter shall be suspended for a period of one year. The license of any provider who has committed three or more violations of this subchapter may be suspended for a period of one year.
(d) Reparation.--Any provider, claimant or other person who is found guilty of a crime for violating this subchapter shall repay three times the value of the material gain received. In addition to the civil penalty authorized pursuant to subsection (b), the department may require the provider, claimant or other person to repay up to three times the value of any material gain to PACE or PACENET.
(521 amended July 7, 2006, P.L.1061, No.111)
Section 522. Prescription drug education program. The department, in cooperation with the Department of Health, shall develop and implement a Statewide prescription drug education program designed to inform older adults of the dangers of prescription drug abuse and misuse. The prescription drug education program shall include, but not be limited to, information concerning the following:
   (1) The hazards of prescription drug overdose.
   (2) The potential dangers of mixing prescription drugs.
   (3) The danger of retaining unused prescription drugs after the need to take them no longer exists.
   (4) The necessity to carefully question physicians, certified registered nurse practitioners and pharmacists concerning the effects of taking prescription drugs,
including the differences between brand-name drugs and generically equivalent drugs.
(5) The advisability of maintaining a prescription drug profile or other record of prescription drug dosage and frequency of dosage.
(6) The desirability of advising family members of the types and proper dosage of prescription drugs which are being taken.
(7) The dangers of taking prescription drugs in excess of prescribed dosages.
(8) The need to obtain complete, detailed directions from the physician, certified registered nurse practitioner or pharmacist concerning the time period a prescription drug should be taken.
(522 amended Nov. 26, 2003, P.L.212, No.37)
Section 522.1. Medication therapy management.
PACE shall, in consultation with the board, develop a proposal for a medication therapy management program by using retail community pharmacies enrolled in the program. PACE, in consultation with the board, shall submit the proposal to the General Assembly no later than six months after the effective date of this section.
(522.1 added Oct. 23, 2018, P.L.579, No.87)

SUBCHAPTER C
COORDINATION OF FEDERAL AND STATE BENEFITS
(Subch. added July 7, 2006, P.L.1045, No.111)

Section 531. Definitions.
The following words and phrases when used in this subchapter shall have the meanings given to them in this section unless the context clearly indicates otherwise:
"LIS." Low-income subsidy assistance from Part D provided by the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (Public Law 108-173, 117 Stat. 2066) to help pay for annual premiums, deductibles and copayments charged to individuals enrolled in Part D by prescription plans approved under that act.
"Part D enrollee." A claimant enrolled in a Part D plan.
"Part D provider." A pharmacy or other prescription drug dispenser authorized by a Part D enrollee's Part D plan.
(531 added July 7, 2006, P.L.1061, No.111)
Section 532. Purpose.
The benefits available to a claimant enrolled in the program under Subchapter B shall be a supplement to the benefits available under Part D. The department may require claimants to utilize Part D benefits prior to utilizing benefits provided under either program and shall coordinate the benefits of the programs with those provided under Part D.
(532 added July 7, 2006, P.L.1061, No.111)
Section 533. Powers of the department.
The department:
(1) Shall identify the Part D plan or plans with which the department has entered into a contract under section 534 that meet the prescription drug needs and pharmacy preferences of a claimant.

(2) May require that the claimant enroll in the Part D plan or program that meets the prescription drug needs and pharmacy preferences of the claimant in the most cost-effective manner for the Commonwealth.

(3) Shall initiate enrollment on behalf of the claimant in the Part D plan recommended by the department unless the claimant notifies the department that the claimant wishes to enroll in another Part D plan.

(4) Shall file and pursue appeals in accordance with CMS regulations with a claimant's Part D plan on the claimant's behalf to request exceptions to the plan's tiered cost-sharing structure or to request a nonformulary Part D drug.

(5) Shall assist claimants the department believes to be eligible for the LIS in making an application to the Social Security Administration.

(6) Shall provide at least ten days for the claimant to decline enrollment in the recommended plan.

(7) Shall develop and distribute language, when recommending enrollment, notifying claimants of:
   (i) The ability to decline enrollment in the recommended Part D plan.
   (ii) The ability to file and pursue appeals to the recommended Part D plan on their own behalf.
   (iii) The possibility that their choice of plan may affect their medical coverage if they are enrolled in a Medicare advantage plan, if applicable.

(533 amended Dec. 22, 2021, P.L.454, No.94)

Section 534. Coordination of benefits.

(a) General coordination.--In addition to the specific provisions of subsection (b), the department shall establish standards and minimum requirements it deems necessary to allow for the coordination of benefits between the program and Part D.

(b) Specific coordination provisions.--The following provisions shall apply to claimants who are also Part D enrollees:

(1) The primary payor shall be the PDP or the Medicare Advantage Prescription Drug Plan, as appropriate.

(2) Part D enrollees shall be required to utilize providers authorized by their PDPs or Medicare Advantage Prescription Drug Plans.

(3) The program shall pay the premium assessed by a PACE enrollee's PDP or, with respect to the prescription drug plan, Medicare Advantage Prescription Drug Plan in an amount not to exceed the regional benchmark premium and any copayments in excess of those set forth in section 509.

(4) Part D enrollees enrolled in PACENET shall pay the Part D premiums charged by their PDP or, with respect to the prescription drug plan, Medicare Advantage Prescription Drug Plan and the program shall pay any copayments in excess of those set forth in section 519. A claimant enrolled in a PDP shall not be required to pay a monthly premium for any month the claimant is not dispensed a prescription drug. (4) amended Dec. 22, 2021, P.L.454, No.94)

(5) For Part D enrollees enrolled in PACE who are not eligible for LIS, PACE shall reimburse Part D providers for prescription drugs in any noncoverage phase of Part D. For
Part D enrollees enrolled in PACENET, PACENET shall reimburse Part D providers for prescription drugs in any noncoverage phase of Part D.

(6) The provisions of Chapter 7 shall apply to all payments made by the program in the noncoverage phase.

(7) The department shall advise a claimant on the various benefits and drugs provided by each PDP approved by the department as follows:

(i) Analyze the claimant's eligibility for and assist the claimant in applying for LIS.

(ii) Identify the claimant's prescription drug needs and preferred pharmacy.

(iii) Assist the claimant in enrolling in the PDP that best fits the claimant's prescription drug needs.

(iv) File and pursue appeals in accordance with CMS regulations with a claimant's Part D plan on the claimant's behalf to request exceptions to the plan's tiered cost-sharing structure or to request a nonformulary Part D drug.

(8) Notwithstanding the provisions of sections 511 and 513(a), for purposes of coordination of benefits with Medicare Part D plans and to minimize disruption to enrollees, the program shall be authorized to reimburse Part D providers, including mail-order pharmacies, for more than a 30-day supply of prescription drugs.

(c) Contracts.--The department is authorized to enter into contracts with Part D plans to provide for prescription drugs to Part D enrollees through Part D pursuant to this subchapter. A Part D plan selected by the department shall meet all of the following requirements:

(1) The Part D plan has a retail pharmacy network that includes at least 90% of the pharmacies in the PACE network.

(2) The Part D plan has a premium at or below the regional benchmark premium.

(c.1) Authorization.--The department may pay the LEP of Part D enrollees in excess of the regional benchmark premium. ((c.1) added Oct. 23, 2018, P.L.579, No.87)

(d) Rebates.--The department may only receive rebates as provided in Chapter 7 where the program is the only payor for a Part D enrollee's covered prescription drugs.

(534 added July 7, 2006, P.L.1061, No.111)

Section 535. Financial resource information.

(a) Procedure.--The department may obtain information on the financial resources of a Part D eligible individual for the purpose of determining the individual's potential eligibility for the LIS. The authority granted under this subsection shall be exercised only with respect to a Part D eligible individual who has income which is below the applicable threshold established by the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (Public Law 108-173, 117 Stat. 2066) for qualification under the LIS.

(b) Waiver.--An application by a Part D eligible individual for enrollment in the program shall constitute a waiver to the department of relevant confidentiality requirements relating to the prospective claimant's financial resources in the possession of any Commonwealth agency or third party when the information is required for the purposes listed under subsection (a). This waiver shall extend to the application phase and throughout the entire time the claimant is in the program.

(c) Information confidential.--

(1) It shall be unlawful for an officer, agent or employee of the department to divulge or make known
information obtained from a Commonwealth agency or third party except for the purposes under subsection (a).

(2) A person that violates this subsection commits a misdemeanor of the third degree and shall, upon conviction, be sentenced to pay a fine of not more than $1,000 or to imprisonment for not more than one year, or both, and to pay the costs of prosecution. If the offender is an officer or employee of the Commonwealth, the offender shall be dismissed from office or discharged from employment.

(d) Upon request of the claimant, the department shall provide a copy of any and all filings that are processed or submitted under this subchapter.

(535 added July 7, 2006, P.L.1061, No.111)

Section 536. Reimbursement.

For-profit insurers, health maintenance organizations, preferred provider organizations, not-for-profit prescription plans, Medicare advantage plans and PDPs shall be responsible for any payments made to a pharmacy on behalf of a Part D enrollee covered by any such third party. Final determination as to the existence of third-party coverage shall be the responsibility of the department.

(536 added July 7, 2006, P.L.1061, No.111)

Section 537. Collection.

The department shall have the authority to collect any amounts from the payment by the department of pharmacy claims that are the responsibility of a PDP or Medicare Advantage Prescription Drug Plan as a primary payor pursuant to section 534(b)(1).

(537 added July 7, 2006, P.L.1061, No.111)

CHAPTER 7

PRUDENT PHARMACEUTICAL PURCHASING

(Hdg. added Nov. 21, 1996, P.L.741, No.134)

Section 701. Declaration of policy.
The General Assembly finds and declares as follows:

(1) The Commonwealth, through assistance programs enacted for the benefit of its citizens, is the largest single payor of prescription medications in Pennsylvania.

(2) In order to ensure that the Commonwealth, in expending money on behalf of its citizens, is not unduly harmed by being required to pay a price for pharmaceutical products purchased from manufacturers in excess of that established for other purchasers and reimbursers of these products and to ensure that the Commonwealth can efficiently and prudently expend its money and maximize its ability to provide for the health and welfare of as many of its needy citizens as possible, it is reasonable, necessary and in the public interest to require that pharmaceutical manufacturers offer a discount to the Commonwealth for pharmaceutical products purchased or reimbursed through State agencies.

(3) It is in the public interest for pharmaceutical manufacturers to provide the Commonwealth with data relating to the price of pharmaceutical products sold by the manufacturer to public bodies, hospitals, for-profit or nonprofit organizations, other manufacturers or wholesalers doing business in this Commonwealth in order to ensure that the Commonwealth can determine that it is being provided with the best prices offered by the manufacturer.

(4) On a national level, there has been a recognition that the need for discounts to State Medicaid agencies, which
reimburse for a high volume of pharmaceutical products, exists.

(5) On a State level, the General Assembly recognizes that it is in the best interest of its citizens to provide pharmaceutical assistance in a reasonable and cost-efficient manner.

(6) Drug price inflation has caused an increase in the amount of public funds expended by PACE and General Assistance.

(701 added Nov. 21, 1996, P.L.741, No.134)

Section 702. Definitions.

The following words and phrases when used in this chapter shall have the meanings given to them in this section unless the context clearly indicates otherwise:

"Average manufacturer price (AMP)." With respect to a covered prescription drug of the manufacturer for a calendar quarter, the average unit price paid to the manufacturer for the drug by wholesalers for drugs distributed to the retail pharmacy class of trade, except for direct sales to hospitals, health maintenance organizations and wholesalers where the drug is relabeled under that distributor's national drug code number. Federal Supply Schedule prices shall not be included in the calculation of AMP. The term includes cash discounts and all other price reductions, other than rebates under this act and section 1927 of Title XIX of the Social Security Act (49 Stat. 620, 42 U.S.C. § 301 et seq.), added November 5, 1990 (Public Law 101-508, Title IV, section 4401(a)(3), 104 Stat. 1388-143), which reduce the actual price paid. For bundled or capitated sales, the allocation of the discount shall be made proportionately to the dollar value of the units of each covered prescription drug sold under the bundled or capitated arrangement. The AMP for a quarter shall be adjusted by the manufacturer if cumulative discounts or other arrangements subsequently adjust the prices actually realized.

"Best price." The lowest price available from the manufacturer during the rebate period to any wholesaler, retailer, provider, health maintenance organization, nonprofit entity or any governmental entity subject to the exclusions and special rules set forth in sections 1902 and 1927(c)(1)(C) of the Social Security Act (49 Stat. 620, 42 U.S.C. §§ 1396c, 1396r-8(c)(1)(C)). (Def. added Nov. 26, 2003, P.L.212, No.37)

"Bundled or capitated sales." The packaging of drugs of different types where:

(1) the condition of rebate or discount is that more than one drug type is purchased; or

(2) the resulting discount or rebate is greater than that which would have been received had the drug products been purchased separately.

"Consumer Price Index-Urban" or "CPI-U." A price index compiled by the Bureau of Labor Statistics of the United States Department of Labor for measuring the average change in the prices paid by urban consumers for a fixed market basket of services.

"Covered prescription drug." A legend drug, insulin, an insulin syringe or an insulin needle eligible for payment by the Commonwealth under PACE, PACENET or designated pharmaceutical programs.

"Depot price." The price available to any depot of the Federal Government for purchase of drugs from the manufacturer through the depot system of procurement.

"Designated pharmaceutical programs." The General Assistance Program and the Special Pharmaceutical Benefit Program in the
Department of Public Welfare and the End Stage Renal Dialysis Program in the Department of Health.

"Direct seller." Any person, partnership, corporation, institution or entity engaged in the selling of pharmaceutical products directly to consumers in this Commonwealth.

"Distributor." A private entity under contract with the original labeler or holder of the national drug code number to manufacture, package or market the covered prescription drug.

"Doing business in this Commonwealth." The direct or indirect selling or the making of covered prescription drugs available for sale in a continuous and systematic manner with the reasonable expectation that these products will be sold to consumers in this Commonwealth.

"FDA." The Food and Drug Administration of the Public Health Service of the Department of Health and Human Services.


"Innovator multiple-source drugs." A multiple-source drug that was originally marketed under a new drug application approved by the FDA. The term includes:

1. covered prescription drugs approved under Product License Approval (PLA), Establishment License Approval (ELA) or Antibiotic Drug Approval (ADA); and
2. a covered prescription drug marketed by a cross-licensed producer or distributor under the approved Abbreviated New Drug Application (ANDA) when the drug product meets this definition.

"Manufacturer." (1) An entity which is engaged in any of the following:
   (i) The production, preparation, propagation, compounding, conversion or processing of prescription drug products:
      (A) directly or indirectly by extraction from substances of natural origin;
      (B) independently by means of chemical synthesis; or
      (C) by a combination of extraction and chemical synthesis.
   (ii) The packaging, repackaging, labeling or relabeling, or distribution of prescription drug products.
(2) The entity holding legal title to or possession of the national drug code number for the covered prescription drug.
(3) The term does not include a wholesale distributor of drugs, drugstore chain organization or retail pharmacy licensed by the Commonwealth.

"National drug code number." The identifying drug number maintained by the FDA. The complete eleven-digit number must include the labeler code, product code and package size code.


"Noninnovator multiple-source drug." Any of the following:
(1) A covered prescription drug which is not an innovator multiple-source drug approved under an Abbreviated New Drug Application (ANDA) or an Amended Antibiotic Drug Approval (AADA).
(2) A drug that has been approved for substitution under the act of November 24, 1976 (P.L.1163, No.259), referred to as the Generic Equivalent Drug Law.

"PACE." The program under Chapter 5.
"PACE." The program established under section 519.
"Private entity." Includes a for-profit entity and a nonprofit entity.
"Producer Price Index for Pharmaceuticals." The prescription
drug producer price index compiled by the Bureau of Labor
Statistics of the United States Department of Labor for
measuring average changes in selling prices received by domestic
drug manufacturers.
"Provider." A licensed pharmacy, dispensing physician or
certified registered nurse practitioner enrolled as a provider
in PACE, PACENET or designated pharmaceutical programs. (Def.
amended Nov. 26, 2003, P.L.212, No.37)
"Rebate period." A calendar quarter or other period
specified by the Secretary of Aging with respect to the payment
of rebates under an agreement as provided in section 703.
"Secretary." The Secretary of Aging of the Commonwealth.
"Single-source drugs." Legend drug products for which the
FDA has not approved an Abbreviated New Drug Application (ANDA).
"Unit." A drug unit in the lowest identifiable amount, such
as tablet or capsule for solid dosage forms, milliliter for
liquid forms and gram for ointments or creams. The manufacturer
shall specify the unit for each dosage form and strength of
each covered prescription drug in accordance with the
instructions developed by the Health Care Financing
Administration for purposes of the Federal Medicaid Rebate
Program under section 1927 of Title XIX of the Social Security
Act (49 Stat. 620, 42 U.S.C. § 301 et seq.).
"Wholesaler." Any person, partnership, corporation,
institution or entity to which the manufacturer sells the
covered prescription drug, including a pharmacy or chain of
pharmacies, but that does not relabel or repackage the covered
prescription drug.
(702 added Nov. 21, 1996, P.L.741, No.134)

Compiler's Note: The Department of Public Welfare, referred
to in this section, was redesignated as the Department
of Human Services by Act 132 of 2014.

Section 703. Rebate agreement.
(a) Requirement.--PACE, PACENET and designated
pharmaceutical programs shall not reimburse for any covered
prescription drug without a rebate agreement between the
department and the manufacturer of the covered prescription
drug.
(b) Exception.--Subsection (a) shall not apply if the
availability of the drug is essential to the health of eligible
claimants as determined by the department.
(c) Agreements.--Manufacturers of prescription drugs
reimbursed under PACE, PACENET and designated pharmaceutical
programs must enter into a rebate agreement with the department
under this chapter to obtain such reimbursement. Nothing in
this chapter shall be deemed to affect or impair any agreement
made under the former provisions of Chapter 6 of the act of
August 14, 1991 (P.L.342, No.36), known as the Lottery Fund
Preservation Act.
(d) Notice.--The department shall notify enrolled providers
of PACE, PACENET and designated pharmaceutical programs on an
annual basis and, as appropriate, of all manufacturers who have
entered into a rebate agreement.
(e) Drug formulary.--Except as provided in section 512,
there shall be no drug formulary, prior or retroactive approval
system or any similar restriction imposed on the coverage of
outpatient drugs made by manufacturers who have agreements in
effect with the Commonwealth to pay rebates for drugs utilized in PACE and PACENET, provided that such outpatient drugs were approved for marketing by the Food and Drug Administration. This subsection shall not apply to any act taken by the department pursuant to its therapeutic drug utilization review program under section 505.

(703 added Nov. 21, 1996, P.L.741, No.134)

Section 704. Terms of rebate agreement.

(a) Quarterly basis.--A rebate agreement shall require any manufacturer of covered prescription drugs to provide to the department a rebate each calendar quarter in an amount specified in section 705 for covered prescription drugs of the manufacturer reimbursed during the quarter. The rebate shall be paid by the manufacturer not later than 30 days after the date of receipt of the information described in subsection (b) for the period involved.

(b) Information.--

(1) The department shall report to each manufacturer, not later than 60 days after the end of each calendar quarter, information by zip code of provider on the total number of dosage units of each covered prescription drug reimbursed under PACE, PACENET and designated pharmaceutical programs during the quarter.

(2) A manufacturer may review the information provided under paragraph (1) and verify information. Adjustments to rebates shall be made to the extent that information indicates that utilization was greater or less than the amount previously specified.

(3) In the event that in any quarter a material discrepancy in the department's information is certified by the manufacturer prior to the due date of the rebate, the department and the manufacturer shall, in good faith, attempt to resolve the discrepancy. If resolution is not reached within 30 days of receipt of the manufacturer's certification by the department, the manufacturer may appeal the department's decision under the department's formal fair hearings and appeals process. The manufacturer shall pay the department that portion of the rebate amount which is not disputed within the required time frame under this chapter. Any balance due, plus statutory interest, shall be paid or credited by the manufacturer or the department by the due date of the next quarterly payment after resolution of the dispute.

(c) Manufacturer provision of price information.--

(1) Each manufacturer with an agreement in effect under this chapter shall report the average manufacturer price and the best price for all covered prescription drugs produced by that manufacturer to the department not later than 30 days after the last day of each quarter. ((1) amended Nov. 26, 2003, P.L.212, No.37)

(2) The department shall retain the services of an independent contractor to survey wholesalers, direct sellers and manufacturers that directly distribute their covered prescription drugs, when necessary, to verify manufacturer prices reported under paragraph (1). Any survey conducted shall not reveal to the department nor to any other person or entity other than the independent contractor the name, identity, location, actual acquisition invoice, other proprietary information or any information from which the department might be enabled to ascertain the name, identity or location of any wholesaler, direct seller or provider so surveyed unless the contractor shall have gathered sufficient
evidence to enable the department to bring charges against any wholesaler, direct seller or provider in violation of subsection (d)(3).

(d) Penalties.--The department shall administer penalties as follows:

(1) A manufacturer who fails to supply information required under subsection (c)(1) shall be liable for a civil penalty in the amount of 2% of the rebate next required to be paid, plus $1,000 for each day that the information is late. If the information is not reported within 30 days of the due date, the agreement shall be suspended for services furnished after the end of the 30-day period until the date the information is reported or the expiration of 45 days, whichever is later.

(2) A manufacturer who knowingly supplies false information that is required under subsection (c)(1) shall be liable for a civil penalty in the amount of $50,000 for each item of false information.

(3) A direct seller, manufacturer or wholesaler who refuses a request for information or knowingly provides false information that is required under subsection (c)(2) shall be liable for a civil penalty in the amount of $50,000.

(4) Penalties collected under this subsection shall be deposited into the fund.

(5) All civil monetary penalties imposed under this chapter are in addition to other civil or criminal penalties.

(e) Confidentiality of information.--Information disclosed by manufacturers, wholesalers or direct sellers under this chapter is confidential and shall not be disclosed by the department in a form which discloses the identity of a specific manufacturer, wholesaler or direct seller or the prices charged for drugs by the manufacturer or wholesaler, except as the department determines to be necessary to carry out this chapter and to permit the Department of the Auditor General and the Office of State Inspector General to review the information provided.

(f) Length of agreement.--A rebate agreement shall remain in effect 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 subsection (g).

(g) Termination.--

(1) The department may provide for termination of a rebate agreement for any reason. Termination shall not be effective earlier than 60 days after the date of receipt of notice of termination by the manufacturers.

(2) A manufacturer may terminate a rebate agreement for any reason. Termination shall not be effective earlier than 60 days after the date of receipt of notice of termination by the department.

(3) Termination of the rebate agreement shall not affect rebates due under the agreement before the effective date of termination.

(4) Commonwealth Court shall have original jurisdiction over cases of termination of agreements under this subsection. Commencement of an action under this paragraph shall not delay the effective date of termination.

(5) If a rebate agreement is terminated for cause, another agreement with the same manufacturer or a successor manufacturer may not be entered into until a period of one year has elapsed from the date of the termination unless the department finds good cause for an earlier agreement.

(704 added Nov. 21, 1996, P.L.741, No.134)
Compiler's Note: Section 15 of Act 37 of 2003, which amended subsection (c)(1), provided that the amendment shall apply retroactively to January 1, 2003.

Section 705. Amount of rebate.

(a) Single-source drugs and innovator multiple-source drugs.--With respect to single-source drugs and innovator multiple-source drugs, each manufacturer shall remit a rebate to the Commonwealth. Except as otherwise provided in this section, the amount of the rebate to the Commonwealth per calendar quarter with respect to each dosage form and strength of single-source drugs and innovator multiple-source drugs shall be as follows:

(1) For quarters beginning after September 30, 1992, and ending before January 1, 1997, the product of the total number of units of each dosage form and strength reimbursed by PACE and General Assistance in the quarter and the difference between the average manufacturer price and 85% of that price, after deducting customary prompt payment discounts, for the quarter.

(2) For quarters beginning after December 31, 1996, and ending before January 1, 2003, the product of the total number of units of each dosage form and strength reimbursed by PACE, PACENET and designated pharmaceutical programs in the quarter and the difference between the average manufacturer price and 83% of that price, after deducting customary prompt payment discounts.

(3) For quarters beginning after December 31, 2002, each manufacturer shall remit a rebate to the Commonwealth for the total number of units of each dosage form and strength reimbursed by PACE, PACENET and designated pharmaceutical programs in the quarter pursuant to the determination established by section 1927(c)(1) of the Social Security Act (49 Stat. 620, 42 U.S.C. § 1396r-8(c)(1)).

(b) Rebate for other drugs.--

(1) The amount of the rebate to the Commonwealth for a calendar quarter with respect to covered prescription drugs which are noninnovator multiple-source drugs shall be equal to the product of:

(i) the applicable percentage of the average manufacturer price, after deducting customary prompt payment discounts, for each dosage form and strength of such drugs for the quarter; and

(ii) the number of units of such form and dosage reimbursed by PACE and General Assistance in the quarter.

(2) For the purposes of paragraph (1), the applicable percentage for calendar quarters beginning after September 30, 1992, and ending before January 1, 1997, is 11%.

(c) Revised rebate for other drugs.--Beginning after December 31, 1996, and ending before January 1, 2004, all of the following shall apply:

(1) The amount of the rebate to the Commonwealth for a calendar quarter with respect to covered prescription drugs which are noninnovator multiple-source drugs shall be the greater of the product of:

(i) The applicable percentage of the average manufacturer price, after deducting customary prompt payment discounts, for each dosage form and strength of such drugs for the quarter; and
the number of units of such form and dosage reimbursed by PACE, PACENET and designated pharmaceutical programs in the quarter.

(2) For purposes of paragraph (1), the applicable percentage is 17%.

(c amended Nov. 26, 2003, P.L.212, No.37)

(c.1) Rebates for other drugs.--For quarters beginning after December 31, 2003, each manufacturer shall remit a rebate to the Commonwealth for the total number of units of each dosage form and strength reimbursed by PACE, PACENET and designated pharmaceutical programs in the quarter pursuant to the determination established by section 1927(c)(3) of the Social Security Act (49 Stat. 620, 42 U.S.C. § 1396r-8(c)(3)). ((c.1) amended Nov. 30, 2004, P.L.1722, No.219)

(d) Drugs approved after act takes effect.--In the case of a covered outpatient drug approved for marketing after the effective date of the act of August 14, 1991 (P.L.342, No.36), known as the Lottery Fund Preservation Act, any reference to January 1, 1991, shall be a reference to the first day of the first month during which the drug was marketed.

(705 added Nov. 21, 1996, P.L.741, No.134)

Section 706. Excessive pharmaceutical price inflation discount.

(a) General rule.--A discount shall be provided to the department for all covered prescription drugs except those excluded under subsection (d). The discount shall be calculated as follows:

(1) For each quarter for which a rebate under section 705(a) and (b) is to be paid after December 31, 1996, the average manufacturer price for each dosage form and strength of a covered prescription drug shall be compared to the average manufacturer price for the same form and strength in the previous calendar year, and a percentage increase shall be calculated.

(2) For each quarter under paragraph (1), the average percentage increase in the Producer Price Index for Pharmaceuticals over the same quarter in the previous calendar year shall be calculated.

(3) If the calculation under paragraph (1) is greater than the calculation under paragraph (2), the discount amount for each quarter shall be equal to the product of:

(i) the difference between the calculations under paragraphs (1) and (2); and

(ii) the total number of units of each dosage form and strength reimbursed by PACE and General Assistance and the average manufacturer price reported by the manufacturer under section 704(c)(1).

(b) Revised general rule.--A discount shall be provided to the department for all covered prescription drugs except those excluded under subsection (d). The discount shall be calculated as follows:

(1) For each quarter for which a rebate under section 705(a) and (c) is to be paid after December 31, 1991, and before January 1, 1997, the average manufacturer price for each dosage form and strength of a covered prescription drug shall be compared to the average manufacturer price for the same form and strength in the previous calendar year, and a percentage increase shall be calculated.

(2) For each quarter under paragraph (1), the average percentage increase in the Consumer Price Index-Urban over the same quarter in the previous calendar year shall be calculated.
If the calculation under paragraph (1) is greater than the calculation under paragraph (2), the discount amount for each quarter shall be equal to the product of:

(i) the difference between the calculations under paragraphs (1) and (2); and
(ii) the total number of units of each dosage form and strength reimbursed by PACE, PACENET and designated pharmaceutical programs and the average manufacturer price reported by the manufacturer under section 704(c)(1).

(c) New bimarked drugs.--For covered prescription drugs that have not been marketed for a full calendar year, subsection (a) shall apply after the covered prescription drug has been on the market for four consecutive quarters. The drug's initial average manufacturer price shall be based on the first day of the first quarter that the drug was marketed.

d) Applicability.--This section shall not apply to a noninnovator multiple-source prescription drug or generic prescription drug.

(706 amended July 7, 2006, P.L.1061, No.111)

Section 707. Lowered best price.

(a) General rule.--If the rebate under section 705 and the discount under section 706 would establish a lowered Federal best price, as defined in section 1927(c)(1)(C) of the Social Security Act (49 Stat. 620, 42 U.S.C. § 1396r-8(c)(1)(C)), the manufacturer shall be liable for a total rebate and discount in an amount that does not reduce the Federal best price for that covered prescription drug.

(b) Procedure.--Any claim by a manufacturer that a rebate would establish a lower Federal best price under subsection (a) shall be verified in writing by a department-approved independent public accounting firm within 45 days of the end of the quarter for which the claim is asserted. The information provided to the public accounting firm shall remain confidential.

(c) Civil penalty.--A manufacturer which provides false information under this section shall be liable for a civil penalty in an amount not to exceed $50,000. Each item of false information constitutes a separate violation.

(707 added Nov. 21, 1996, P.L.741, No.134)

Section 708. Exemption.

Section 521(a) shall not apply to rebates under this chapter.

(708 added Nov. 21, 1996, P.L.741, No.134)

Section 709. Disposition of funds.

(a) PACE and PACENET.--Money received under this chapter in connection with PACE and PACENET shall be deposited in the Pharmaceutical Assistance Contract for the Elderly Fund.

(b) Designated pharmaceutical programs.--Money received under this chapter in connection with designated pharmaceutical programs shall be treated as a refund of expenditures to the appropriation which originally provided the funding for the pharmaceutical purchase.

(709 added Nov. 21, 1996, P.L.741, No.134)

CHAPTER 8

PHARMACEUTICAL ASSISTANCE CLEARINGHOUSE

(Hdg. added Nov. 26, 2003, P.L.212, No.37)
"Clearinghouse." The Pharmaceutical Assistance Clearinghouse established in section 802.

"Department." The Department of Aging of the Commonwealth.

"Patient assistance program." A program offered by a pharmaceutical manufacturer under which the manufacturer provides prescription medications at no charge or at a substantially reduced cost. The term does not include the provision of a drug as part of a clinical trial.

"Voluntary health organization." An organization whose main purpose is to educate the public on the symptoms, treatments and research of a disease and that may provide support for persons who have the disease.

(801 added Nov. 26, 2003, P.L.212, No.37)

Section 802. Pharmaceutical Assistance Clearinghouse.

(a) Establishment.--Within 120 days of the effective date of this chapter, the department shall establish the Pharmaceutical Assistance Clearinghouse. Each pharmaceutical manufacturer that does business in this Commonwealth and offers a patient assistance program shall inform the department of all of the following:

(1) The existence of the patient assistance program.
(2) The eligibility requirements for the patient assistance program.
(3) The drugs covered by the patient assistance program.
(4) Information, such as a telephone number, which may be used to apply for a patient assistance program.

(b) Information.--The clearinghouse shall maintain the information submitted by pharmaceutical manufacturers and any appropriate voluntary health organization that would like to participate and make it available to the public.

(c) Staff.--The department shall ensure that the clearinghouse is staffed at least during normal business hours.

(802 added Nov. 26, 2003, P.L.212, No.37)

Section 803. Toll-free telephone number.
The department shall establish a toll-free telephone number through which members of the public may obtain information from the clearinghouse about available patient assistance programs.

(803 added Nov. 26, 2003, P.L.212, No.37)

Section 804. Assistance available.

(a) Direct.--

(1) The clearinghouse shall assist without charge an individual in determining whether a patient assistance program is offered for a particular drug and whether the individual may be eligible to obtain the drug through a patient assistance program.

(2) The clearinghouse may assist without charge an individual who wishes to apply for a patient assistance program by assisting with the preparation of an application and coordinating communications between the individual's physician or certified registered nurse practitioner and a pharmaceutical manufacturer on behalf of the individual for the purpose of obtaining approval to participate in the patient assistance program.

(b) Referrals.--The clearinghouse shall make referrals to an appropriate voluntary health organization or any publicly funded program for which it deems a patient eligible.

(804 added Nov. 26, 2003, P.L.212, No.37)

Section 805. Reporting.
The department shall report annually to the Governor and the General Assembly on the activities of the clearinghouse. The report shall include:
(1) The number of individuals who have been assisted by the clearinghouse under section 804(a)(1) and the number of such individuals under section 804(a)(2).

(2) The number and benefits of patient assistance programs listed with the clearinghouse.

(3) The number of patients referred to publicly funded programs under section 804(b). Programs under this paragraph include, but are not limited to, the Pharmaceutical Assistance Contract for the Elderly program, medical assistance and programs of the Department of Veterans Affairs.

(4) Other information deemed relevant by the department.

(805 added Nov. 26, 2003, P.L.212, No.37)

Section 806. Internet availability of information.
The department shall maintain and provide to the public the information under this chapter on its World Wide Web site. The department shall also provide to appropriate organizations the information necessary for the organizations to establish a link to the location of clearinghouse information on the department's World Wide Web site.

(806 added Nov. 26, 2003, P.L.212, No.37)

CHAPTER 9
TRANSPORTATION SERVICES
(Hdg. added Nov. 21, 1996, P.L.741, No.134)

Compiler's Note: Section 2 of Act 30 of 1999 provided that as much of Chapter 9 as relates to the payment of administrative expenses, other than the payment of commissions, of Act 91 is repealed.

Section 901. Definitions.
The following words and phrases when used in this chapter shall have the meanings given to them in this section unless the context clearly indicates otherwise:

"Shared-ride public transportation services."
Demand-responsive transportation that is available to the general public, operates on a nonfixed route basis and charges a fare to all riders. For transportation to be included in this definition, the first fare-paying passengers to enter the public transportation vehicle must not refuse to share the vehicle with other passengers during a given trip. The term excludes exclusive ride taxi service, charter and sightseeing services, nonpublic transportation, school bus and limousine services.

(901 added Nov. 21, 1996, P.L.741, No.134)

Section 902. Department of Transportation.
The Department of Transportation has the power and duty to make grants from the fund to transportation companies, county transportation systems and local transportation organizations to pay estimated transit losses resulting from providing free service or local common carrier mass transportation systems to persons 65 years of age or older when passage is on fixed-route public transportation services during nonpeak riding hours and on holidays and weekends. Reimbursement shall be as follows:

(1) The losses resulting from granting service on mass transportation systems shall be reimbursable at 100% of the system's average or base fare, whichever is less, multiplied by the number of trips made by senior citizens participating in the free transit program.

(2) Notwithstanding paragraph (1), the department shall, with the approval of the Governor's Office of the Budget, reimburse transportation companies or local transportation
organizations for 100% of the losses resulting from senior citizen transfer trips incurred under the conditions of this subsection.

(3) Money appropriated from the General Fund to the Department of Transportation to augment fixed-route public transportation services under this section shall be granted to transportation providers at the discretion of the Department of Transportation.

(902 added Nov. 21, 1996, P.L.741, No.134)

Section 903. Commuter rail fare.

With regard to passage on fixed-route commuter rail lines, the fare for adults 65 years of age or older who do not qualify as eligible claimants under the pharmaceutical assistance contract for the elderly program shall be limited to $1 per trip but only when utilizing such transportation services during nonpeak riding hours and on holidays and weekends.

(903 added Nov. 21, 1996, P.L.741, No.134)

Section 904. Shared-ride transportation.

(a) Program grants.--The Department of Transportation has the power and duty to administer, utilizing a fixed amount of money from the fund as provided through executive authorizations by the Governor, a program providing shared-ride public transportation services for adults 65 years of age or older. Individuals utilizing shared-ride public transportation services for older adults shall contribute 15% of the individual fare and 85% of the individual fare shall be reimbursed by the fund.

(b) Regulations.--The Department of Transportation shall promulgate regulations necessary to carry out the purposes of this section, including regulations that permit limited reimbursement for shared-ride public transportation services providing access to and from public airports. The Department of Transportation, in consultation with the Pennsylvania Public Utility Commission and the department, shall establish reasonable per mile or trip fare limits for purposes of subsection (a). In accordance with section 2203-A(a)(27) of the act of April 9, 1929 (P.L.177, No.175), known as The Administrative Code of 1929, no regulation shall take effect until submitted to the department for comment.

(c) Coordinated transportation plans.--The Department of Transportation shall require that each transportation provider or designated coordinator, whichever is appropriate, annually develop a coordinated transportation plan which shall include, but not be limited to, any current fixed-route system and shared-ride programs. All plans must be submitted to and approved by the department no later than June 1 of each fiscal year.

(d) Other forms of assistance.--Any eligible claimant whose transportation services are covered in part by any other plan of assistance may be required to receive reduced transportation assistance under the provisions of this chapter.

(e) Entitlement not created.--Nothing in this chapter creates or provides any individual with an entitlement to services.

(904 added Nov. 21, 1996, P.L.741, No.134)

Section 905. Grants.

Grants may be made under this chapter with reference to any appropriate project, irrespective of when it was first commenced or considered and regardless of whether costs with respect to the project have been incurred prior to the time the grant is applied for or made.

(905 added Nov. 21, 1996, P.L.741, No. 134)
Section 2101. Savings.
This act does not affect any act done, regulation promulgated, liability incurred or right accrued or vested or affect any civil or criminal proceeding pending or to be commenced to enforce any right or penalty or punish any offense under any statute or part of a statute repealed by this act.
(2101 added Nov. 21, 1996, P.L.741, No.134)

Section 2102. Annual report to General Assembly.
(a) Submission of report.--The department shall submit a report no later than April 1 of each year to the chairman and minority chairman of the Aging and Youth Committee of the Senate, the chairman and minority chairman of the Aging and Older Adult Services Committee of the House of Representatives and the Pharmaceutical Assistance Review Board. ((a) amended Nov. 26, 2003, P.L.212, No.37)
(b) Collection of data.--The department shall maintain monthly statistical records on PACE and PACENET, including the level of participation and any patterns of unusual drug usage for purposes of formulating the annual report.
(c) Information for inclusion in annual report.--The annual report shall contain, but not be limited to, all information relating to:
   (1) The number of persons served by PACE and PACENET and their counties of residence.
   (2) A breakdown of the numbers and kinds of pharmaceuticals used.
   (3) The cost of prescriptions.
   (4) An estimate of actual expenses incurred by pharmacists participating in the program.
   (5) The results obtained by the drug education program under section 522.
   (6) Information regarding the operation of the therapeutic drug utilization review system for the prior calendar year, which shall include, at a minimum:
      (i) The scope of physician and pharmacist participation in the system.
      (ii) A description of claimant response to the system.
      (iii) Data for each month of the covered period regarding the number of prescription revisions based on utilization review, including drug information, cost savings and the policy used by the department to make utilization review decisions.
   (7) Information on the existence and scope of fraudulent activity and violations of this act by providers participating in PACE and PACENET.
   (8) Information regarding the financial status of PACE and PACENET, including, but not limited to, the adequacy of any applicable deductible and copayment levels, based upon the financial experience and projections of PACE and PACENET.
(2102 added Nov. 21, 1996, P.L.741, No.134)

Section 2103. Federal programs.
If the Federal Government enacts pharmacy programs similar to PACE or PACENET, the State programs shall be construed to only supplement the Federal pharmacy programs. All persons qualified for coverage under a Federal pharmacy program, including the prescription drug benefit program provided by the Medicare Prescription Drug, Improvement, and Modernization Act
of 2003 (Public Law 108-173, 117 Stat. 2066), may be required by the department to utilize the Federal program before utilizing any State program. 

(2103 amended July 7, 2006, P.L.1061, No.111)

APPENDIX

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Supplementary Provisions of Amendatory Statutes
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2006, JULY 7, P.L.1061, NO.111

Section 14. (a) Notwithstanding any other provision of law to the contrary, persons who, as of December 31, 2005, are enrolled in the PACE or PACENET program as defined in section 502 of the act shall remain eligible for the PACE or PACENET program if the maximum income limit is exceeded due solely to a Social Security cost-of-living adjustment.

(b) Funding, to the extent authorized by section 306(b)(1)(vii) of the act of June 26, 2001 (P.L.755, No.77), known as the Tobacco Settlement Act, shall continue to be appropriated to the Pharmaceutical Assistance Contract for the Elderly Fund to support the program expansions contained in this section. The Department of Aging shall also designate funds from the fund to continue eligibility under this section; however, these funds shall not exceed the funding designated under section 306(b)(1)(vii) of the Tobacco Settlement Act. If eligibility under this section requires that funds from the fund exceed those from section 306(b)(1)(vii) of the Tobacco Settlement Act, then the Department of Aging is authorized to determine eligibility requirements.

(c) Eligibility in the PACE program pursuant to this section shall expire December 31, 2006.

(d) Eligibility in the PACENET program pursuant to this section shall expire December 31, 2007.

Compiler's Note: Act 111 amended, added or deleted the heading of Subchapter A of Chapter 5, section 502, the heading of Subchapter 518, 519, er 5 and sections 504, 505, 506, 509, 510, 512, 515, 517, 520.1, and 521, the Subchapter C heading of Chapter 5 and sections 531, 532, 533, 534, 535, 536, 537, 706 and 2103 of Act 91.