A bill to be entitled

An act relating to Medicaid; amending s. 409.904, F.S.; providing for funding the Medicaid reimbursement for certain persons age 65 or older while the optional program is being phased out; renaming the "medically needy" program as the "Medicaid nonpoverty medical subsidy"; limiting certain categories of persons eligible for the subsidy to only physician services after a certain date; amending s. 409.905, F.S.; deleting the hospitalist program; amending s. 409.908, F.S.; revising the factors for calculating the maximum allowable fee for pharmaceutical ingredient costs; directing the Agency for Health Care Administration to establish reimbursement rates for the next fiscal year; amending s. 409.9082, F.S.; revising the aggregated amount of the quality assessment for nursing home facilities; exempting certain nursing home facilities from the quality assessment; amending s. 409.911, F.S.; updating references to data to be used for the disproportionate share program; amending s. 409.9112, F.S.; extending the prohibition against distributing moneys under the regional perinatal intensive care centers disproportionate share program for another year; amending s. 409.9113, F.S.; extending the disproportionate share program for teaching hospitals for another year; amending s. 409.9117, F.S.; extending the prohibition against distributing moneys under the primary care disproportionate share program for another year;
amending s. 409.912, F.S.; allowing the agency to
continue to contract for electronic access to certain
pharmacology drug information; eliminating the
requirement to implement a wireless handheld clinical
pharmacology drug information database for
practitioners; revising the factors for calculating
the maximum allowable fee for pharmaceutical
ingredient costs; amending ss. 409.9122, 409.915, and
409.9301, F.S.; conforming provisions to changes made
by the act; providing an effective date.

Be It Enacted by the Legislature of the State of Florida:

Section 1. Subsections (1) and (2) of section 409.904,
Florida Statutes, are amended to read:

409.904 Optional payments for eligible persons.—The agency
may make payments for medical assistance and related services on
behalf of the following persons who are determined to be
eligible subject to the income, assets, and categorical
eligibility tests set forth in federal and state law. Payment on
behalf of these Medicaid eligible persons is subject to the
availability of moneys and any limitations established by the
General Appropriations Act or chapter 216.

(1) Effective January 1, 2006, and Subject to federal
waiver approval, a person who is age 65 or older or is
determined to be disabled, whose income is at or below 88
percent of the federal poverty level, whose assets do not exceed
established limitations, and who is not eligible for Medicare
or, if eligible for Medicare, is also eligible for and receiving
Medicaid-covered institutional care services, hospice services, or home and community-based services. The agency shall seek federal authorization through a waiver to provide this coverage. This eligibility category subsection expires June 30, 2011. However, for the purpose of phasing out this category, the agency may continue making payments through March 31, 2012.

(2) (a) A family, a pregnant woman, a child under age 21, a person age 65 or over, or a blind or disabled person, who would be eligible under any group listed in s. 409.903(1), (2), or (3), except that the income or assets of such family or person exceed established limitations is eligible for the Medicaid nonpoverty medical subsidy, which includes the same services as those provided to other Medicaid recipients, with the exception of services in skilled nursing facilities and intermediate care facilities for the developmentally disabled. For a family or person in one of these coverage groups, medical expenses are deductible from income in accordance with federal requirements in order to make a determination of eligibility. Effective April 1, 2012, a family, a person age 65 or older, or a blind or disabled person is eligible to receive physician services only. A family or person eligible under the coverage known as the “medically needy,” is eligible to receive the same services as other Medicaid recipients, with the exception of services in skilled nursing facilities and intermediate care facilities for the developmentally disabled. This paragraph expires June 30, 2011.

(b) Effective July 1, 2011, a pregnant woman or a child younger than 21 years of age who would be eligible under any group listed in s. 409.903, except that the income or assets of
such group exceed established limitations. For a person in one of these coverage groups, medical expenses are deductible from income in accordance with federal requirements in order to make a determination of eligibility. A person eligible under the coverage known as the “medically needy” is eligible to receive the same services as other Medicaid recipients, with the exception of services in skilled nursing facilities and intermediate care facilities for the developmentally disabled.

Section 2. Paragraphs (d), (e), and (f) of subsection (5) of section 409.905, Florida Statutes, are amended to read:

409.905 Mandatory Medicaid services.—The agency may make payments for the following services, which are required of the state by Title XIX of the Social Security Act, furnished by Medicaid providers to recipients who are determined to be eligible on the dates on which the services were provided. Any service under this section shall be provided only when medically necessary and in accordance with state and federal law. Mandatory services rendered by providers in mobile units to Medicaid recipients may be restricted by the agency. Nothing in this section shall be construed to prevent or limit the agency from adjusting fees, reimbursement rates, lengths of stay, number of visits, number of services, or any other adjustments necessary to comply with the availability of moneys and any limitations or directions provided for in the General Appropriations Act or chapter 216.

(5) HOSPITAL INPATIENT SERVICES.—The agency shall pay for all covered services provided for the medical care and treatment of a recipient who is admitted as an inpatient by a licensed physician or dentist to a hospital licensed under part I of
chapter 395. However, the agency shall limit the payment for inpatient hospital services for a Medicaid recipient 21 years of age or older to 45 days or the number of days necessary to comply with the General Appropriations Act.

(d) The agency shall implement a hospitalist program in nonteaching hospitals, select counties, or statewide. The program shall require hospitalists to manage Medicaid recipients’ hospital admissions and lengths of stay. Individuals who are dually eligible for Medicare and Medicaid are exempted from this requirement. Medicaid participating physicians and other practitioners with hospital admitting privileges shall coordinate and review admissions of Medicaid recipients with the hospitalist. The agency may competitively bid a contract for selection of a single qualified organization to provide hospitalist services. The agency may procure hospitalist services by individual county or may combine counties in a single procurement. The qualified organization shall contract with or employ board-eligible physicians in Miami-Dade, Palm Beach, Hillsborough, Pasco, and Pinellas Counties. The agency is authorized to seek federal waivers to implement this program.

(e) The agency shall implement a comprehensive utilization management program for hospital neonatal intensive care stays in certain high-volume participating hospitals, select counties, or statewide, and shall replace existing hospital inpatient utilization management programs for neonatal intensive care admissions. The program shall be designed to manage the lengths of stay for children being treated in neonatal intensive care units and must seek the earliest medically appropriate discharge to the child’s home or other
less costly treatment setting. The agency may competitively bid a contract for the selection of a qualified organization to provide neonatal intensive care utilization management services. The agency may be authorized to seek any federal waivers to implement this initiative.

(e) (f) The agency may develop and implement a program to reduce the number of hospital readmissions among the non-Medicare population eligible in areas 9, 10, and 11.

Section 3. Subsections (14) and (23) of section 409.908, Florida Statutes, are amended to read:

409.908 Reimbursement of Medicaid providers.—Subject to specific appropriations, the agency shall reimburse Medicaid providers, in accordance with state and federal law, according to methodologies set forth in the rules of the agency and in policy manuals and handbooks incorporated by reference therein. These methodologies may include fee schedules, reimbursement methods based on cost reporting, negotiated fees, competitive bidding pursuant to s. 287.057, and other mechanisms the agency considers efficient and effective for purchasing services or goods on behalf of recipients. If a provider is reimbursed based on cost reporting and submits a cost report late and that cost report would have been used to set a lower reimbursement rate for a rate semester, then the provider’s rate for that semester shall be retroactively calculated using the new cost report, and full payment at the recalculated rate shall be effected retroactively. Medicare-granted extensions for filing cost reports, if applicable, shall also apply to Medicaid cost reports. Payment for Medicaid compensable services made on behalf of Medicaid eligible persons is subject to the
availability of moneys and any limitations or directions provided for in the General Appropriations Act or chapter 216. Further, nothing in this section shall be construed to prevent or limit the agency from adjusting fees, reimbursement rates, lengths of stay, number of visits, or number of services, or making any other adjustments necessary to comply with the availability of moneys and any limitations or directions provided for in the General Appropriations Act, provided the adjustment is consistent with legislative intent.

(14) A provider of prescribed drugs shall be reimbursed the least of the amount billed by the provider, the provider’s usual and customary charge, or the Medicaid maximum allowable fee established by the agency, plus a dispensing fee. The Medicaid maximum allowable fee for ingredient cost must will be based on the lowest lower of: the average wholesale price (AWP) minus 16.4 percent, the wholesaler acquisition cost (WAC) plus 1.5 4.75 percent, the federal upper limit (FUL), the state maximum allowable cost (SMAC), or the usual and customary (UAC) charge billed by the provider.

(a) Medicaid providers must are required to dispense generic drugs if available at lower cost and the agency has not determined that the branded product is more cost-effective, unless the prescriber has requested and received approval to require the branded product.

(b) The agency shall is directed to implement a variable dispensing fee for payments for prescribed medicines while ensuring continued access for Medicaid recipients. The variable dispensing fee may be based upon, but not limited to, either or both the volume of prescriptions dispensed by a specific
pharmacy provider, the volume of prescriptions dispensed to an individual recipient, and dispensing of preferred-drug-list products.

(c) The agency may increase the pharmacy dispensing fee authorized by statute and in the annual General Appropriations Act by $0.50 for the dispensing of a Medicaid preferred-drug-list product and reduce the pharmacy dispensing fee by $0.50 for the dispensing of a Medicaid product that is not included on the preferred drug list.

(d) The agency may establish a supplemental pharmaceutical dispensing fee to be paid to providers returning unused unit-dose packaged medications to stock and crediting the Medicaid program for the ingredient cost of those medications if the ingredient costs to be credited exceed the value of the supplemental dispensing fee.

(e) The agency may be authorized to limit reimbursement for prescribed medicine in order to comply with any limitations or directions provided for in the General Appropriations Act, which may include implementing a prospective or concurrent utilization review program.

(23) (a) The agency shall establish rates at a level that ensures no increase in statewide expenditures resulting from a change in unit costs for 2 fiscal years effective July 1, 2009.

(a) Reimbursement rates for the 2011-2012 state fiscal year 2 fiscal years shall be as provided in the General Appropriations Act.

(b) This subsection applies to the following provider types:

1. Inpatient hospitals.
2. Outpatient hospitals.
3. Nursing homes.
4. County health departments.
5. Community intermediate care facilities for the
developmentally disabled.
6. Prepaid health plans.

(c) The agency shall apply the effect of this subsection to
the reimbursement rates for nursing home diversion programs.

(c) The agency shall create a workgroup on hospital
reimbursement, a workgroup on nursing facility reimbursement,
and a workgroup on managed care plan payment. The workgroups
shall evaluate alternative reimbursement and payment
methodologies for hospitals, nursing facilities, and managed
care plans, including prospective payment methodologies for
hospitals and nursing facilities. The nursing facility workgroup
shall also consider price-based methodologies for indirect care
and acuity adjustments for direct care. The agency shall submit
a report on the evaluated alternative reimbursement
methodologies to the relevant committees of the Senate and the
House of Representatives by November 1, 2009.

(d) This subsection expires June 30, 2012.

Section 4. Subsection (2) and paragraph (d) of subsection
(3) of section 409.9082, Florida Statutes, are amended to read:

409.9082 Quality assessment on nursing home facility
providers; exemptions; purpose; federal approval required;
remedies.—

(2) Effective April 1, 2009, a quality assessment there is
imposed upon each nursing home facility a quality assessment.
The aggregated amount of assessments for all nursing home
facilities in a given year may be an amount not exceed the maximum percentage exceeding 5.5 percent of the total aggregate net patient service revenue of assessed facilities allowed under federal law. The agency shall calculate the quality assessment rate annually on a per-resident-day basis, exclusive of those resident days funded by the Medicare program, as reported by the facilities. The per-resident-day assessment rate must be uniform except as prescribed in subsection (3). Each facility shall report monthly to the agency its total number of resident days, exclusive of Medicare Part A resident days, and shall remit an amount equal to the assessment rate times the reported number of days. The agency shall collect, and each facility shall pay, the quality assessment each month. The agency shall collect the assessment from nursing home facility providers by no later than the 15th day of the next succeeding calendar month. The agency shall notify providers of the quality assessment and provide a standardized form to complete and submit with payments. The collection of the nursing home facility quality assessment shall commence no sooner than 5 days after the agency’s initial payment of the Medicaid rates containing the elements prescribed in subsection (4). Nursing home facilities may not create a separate line-item charge for the purpose of passing through the assessment through to residents.

(3)

(d) Effective July 1, 2011 2009, the agency may exempt from the quality assessment any or apply a lower quality assessment rate to a qualified public, nonstate-owned or operated nursing home facility whose total annual indigent
census days are greater than \( \frac{15}{25} \) percent of the facility's total annual census days.

Section 5. Paragraph (a) of subsection (2) of section 409.911, Florida Statutes, is amended to read:

409.911 Disproportionate share program.—Subject to specific allocations established within the General Appropriations Act and any limitations established pursuant to chapter 216, the agency shall distribute, pursuant to this section, moneys to hospitals providing a disproportionate share of Medicaid or charity care services by making quarterly Medicaid payments as required. Notwithstanding the provisions of s. 409.915, counties are exempt from contributing toward the cost of this special reimbursement for hospitals serving a disproportionate share of low-income patients.

(2) The Agency for Health Care Administration shall use the following actual audited data to determine the Medicaid days and charity care to be used in calculating the disproportionate share payment:


Section 6. Section 409.9112, Florida Statutes, is amended to read:

409.9112 Disproportionate share program for regional perinatal intensive care centers.—In addition to the payments made under s. 409.911, the agency shall design and implement a system for making disproportionate share payments to those hospitals that participate in the regional perinatal intensive
care center program established pursuant to chapter 383. The system of payments must conform to federal requirements and distribute funds in each fiscal year for which an appropriation is made by making quarterly Medicaid payments. Notwithstanding s. 409.915, counties are exempt from contributing toward the cost of this special reimbursement for hospitals serving a disproportionate share of low-income patients. For the 2011-2012 state fiscal year, the agency may not distribute moneys under the regional perinatal intensive care centers disproportionate share program.

(1) The following formula shall be used by the agency to calculate the total amount earned for hospitals that participate in the regional perinatal intensive care center program:

\[
TAE = \frac{HDSP}{THDSP}
\]

Where:

- **TAE** = total amount earned by a regional perinatal intensive care center.
- **HDSP** = the prior state fiscal year regional perinatal intensive care center disproportionate share payment to the individual hospital.
- **THDSP** = the prior state fiscal year total regional perinatal intensive care center disproportionate share payments to all hospitals.

(2) The total additional payment for hospitals that participate in the regional perinatal intensive care center program shall be calculated by the agency as follows:
TAP = TAE x TA

Where:

TAP = total additional payment for a regional perinatal intensive care center.

TAE = total amount earned by a regional perinatal intensive care center.

TA = total appropriation for the regional perinatal intensive care center disproportionate share program.

(3) In order to receive payments under this section, a hospital must be participating in the regional perinatal intensive care center program pursuant to chapter 383 and must meet the following additional requirements:

(a) Agree to conform to all departmental and agency requirements to ensure high quality in the provision of services, including criteria adopted by departmental and agency rule concerning staffing ratios, medical records, standards of care, equipment, space, and such other standards and criteria as the department and agency deem appropriate as specified by rule.

(b) Agree to provide information to the Department of Health and the agency, in a form and manner to be prescribed by rule of the department and agency, concerning the care provided to all patients in neonatal intensive care centers and high-risk maternity care.

(c) Agree to accept all patients for neonatal intensive care and high-risk maternity care, regardless of ability to pay, on a functional space-available basis.
(d) Agree to develop arrangements with other maternity and neonatal care providers in the hospital’s region for the appropriate receipt and transfer of patients in need of specialized maternity and neonatal intensive care services.

(e) Agree to establish and provide a developmental evaluation and services program for certain high-risk neonates, as prescribed and defined by rule of the department.

(f) Agree to sponsor a program of continuing education in perinatal care for health care professionals within the region of the hospital, as specified by rule.

(g) Agree to provide backup and referral services to the county health departments and other low-income perinatal providers within the hospital’s region, including the development of written agreements between these organizations and the hospital.

(h) Agree to arrange for transportation for high-risk obstetrical patients and neonates in need of transfer from the community to the hospital or from the hospital to another more appropriate facility.

(4) Hospitals that fail to comply with any of the conditions in subsection (3) or the applicable rules of the Department of Health and the agency may not receive any payments under this section until full compliance is achieved. A hospital that is not in compliance in two or more consecutive quarters may not receive its share of the funds. Any forfeited funds shall be distributed by the remaining participating regional perinatal intensive care center program hospitals.

Section 7. Section 409.9113, Florida Statutes, is amended to read:
409.9113 Disproportionate share program for teaching hospitals.—In addition to the payments made under ss. 409.911 and 409.9112, the agency shall make disproportionate share payments to statutorily defined teaching hospitals, as defined in s. 408.07, for their increased costs associated with medical education programs and for tertiary health care services provided to the indigent. This system of payments must conform to federal requirements and distribute funds in each fiscal year for which an appropriation is made by making quarterly Medicaid payments. Notwithstanding s. 409.915, counties are exempt from contributing toward the cost of this special reimbursement for hospitals serving a disproportionate share of low-income patients. For the 2011-2012 state fiscal year, the agency shall distribute the moneys provided in the General Appropriations Act to statutorily defined teaching hospitals and family practice teaching hospitals, as defined in s. 395.805, pursuant to this section under the teaching hospital disproportionate share program. The funds provided for statutorily defined teaching hospitals shall be distributed in the same proportion as the state fiscal year 2003-2004 state fiscal year teaching hospital disproportionate share funds were distributed or as otherwise provided in the General Appropriations Act. The funds provided for family practice teaching hospitals shall be distributed equally among family practice teaching hospitals.

(1) On or before September 15 of each year, the agency shall calculate an allocation fraction to be used for distributing funds to state statutory teaching hospitals. Subsequent to the end of each quarter of the state fiscal year,
the agency shall distribute to each statutory teaching hospital, as defined in s. 408.07, an amount determined by multiplying one-fourth of the funds appropriated for this purpose by the Legislature times such hospital’s allocation fraction. The allocation fraction for each such hospital shall be determined by the sum of the following three primary factors, divided by three:

(a) The number of nationally accredited graduate medical education programs offered by the hospital, including programs accredited by the Accreditation Council for Graduate Medical Education and the combined Internal Medicine and Pediatrics programs acceptable to both the American Board of Internal Medicine and the American Board of Pediatrics at the beginning of the state fiscal year preceding the date on which the allocation fraction is calculated. The numerical value of this factor is the fraction that the hospital represents of the total number of programs, where the total is computed for all state statutory teaching hospitals.

(b) The number of full-time equivalent trainees in the hospital, which comprises two components:

1. The number of trainees enrolled in nationally accredited graduate medical education programs, as defined in paragraph (a). Full-time equivalents are computed using the fraction of the year during which each trainee is primarily assigned to the given institution, over the state fiscal year preceding the date on which the allocation fraction is calculated. The numerical value of this factor is the fraction that the hospital represents of the total number of full-time equivalent trainees enrolled in accredited graduate programs, where the total is
2. The number of medical students enrolled in accredited colleges of medicine and engaged in clinical activities, including required clinical clerkships and clinical electives. Full-time equivalents are computed using the fraction of the year during which each trainee is primarily assigned to the given institution, over the course of the state fiscal year preceding the date on which the allocation fraction is calculated. The numerical value of this factor is the fraction that the given hospital represents of the total number of full-time equivalent students enrolled in accredited colleges of medicine, where the total is computed for all state statutory teaching hospitals.

The primary factor for full-time equivalent trainees is computed as the sum of these two components, divided by two.

(c) A service index that comprises three components:

1. The Agency for Health Care Administration Service Index, computed by applying the standard Service Inventory Scores established by the agency to services offered by the given hospital, as reported on Worksheet A-2 for the last fiscal year reported to the agency before the date on which the allocation fraction is calculated. The numerical value of this factor is the fraction that the given hospital represents of the total Agency for Health Care Administration Service index values, where the total is computed for all state statutory teaching hospitals.

2. A volume-weighted service index, computed by applying the standard Service Inventory Scores established by the agency
for Health Care Administration to the volume of each service, expressed in terms of the standard units of measure reported on Worksheet A-2 for the last fiscal year reported to the agency before the date on which the allocation factor is calculated. The numerical value of this factor is the fraction that the given hospital represents of the total volume-weighted service index values, where the total is computed for all state statutory teaching hospitals.

3. Total Medicaid payments to each hospital for direct inpatient and outpatient services during the fiscal year preceding the date on which the allocation factor is calculated. This includes payments made to each hospital for such services by Medicaid prepaid health plans, whether the plan was administered by the hospital or not. The numerical value of this factor is the fraction that each hospital represents of the total of such Medicaid payments, where the total is computed for all state statutory teaching hospitals.

The primary factor for the service index is computed as the sum of these three components, divided by three.

(2) By October 1 of each year, the agency shall use the following formula to calculate the maximum additional disproportionate share payment for statutory statutorily defined teaching hospitals:

\[ TAP = \text{THAF} \times A \]

Where:

TAP = total additional payment.
THAF = teaching hospital allocation factor.
A = amount appropriated for a teaching hospital disproportionate share program.

Section 8. Section 409.9117, Florida Statutes, is amended to read:

409.9117 Primary care disproportionate share program.—For the 2011-2012 state fiscal year, the agency may shall not distribute moneys under the primary care disproportionate share program.

(1) If federal funds are available for disproportionate share programs in addition to those otherwise provided by law, there shall be created a primary care disproportionate share program shall be established.

(2) The following formula shall be used by the agency to calculate the total amount earned for hospitals that participate in the primary care disproportionate share program:

\[ TAE = \frac{HDSP}{THDSP} \]

Where:

TAE = total amount earned by a hospital participating in the primary care disproportionate share program.

HDSP = the prior state fiscal year primary care disproportionate share payment to the individual hospital.

THDSP = the prior state fiscal year total primary care disproportionate share payments to all hospitals.

(3) The total additional payment for hospitals that participate in the primary care disproportionate share program

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shall be calculated by the agency as follows:

\[ TAP = TAE \times TA \]

Where:

- **TAP** = total additional payment for a primary care hospital.
- **TAE** = total amount earned by a primary care hospital.
- **TA** = total appropriation for the primary care disproportionate share program.

(4) In establishing the establishment and funding of this program, the agency shall use the following criteria in addition to those specified in s. 409.911, and payments may not be made to a hospital unless the hospital agrees to:

   (a) Cooperate with a Medicaid prepaid health plan, if one exists in the community.

   (b) Ensure the availability of primary and specialty care physicians to Medicaid recipients who are not enrolled in a prepaid capitated arrangement and who are in need of access to such physicians.

   (c) Coordinate and provide primary care services free of charge, except copayments, to all persons with incomes up to 100 percent of the federal poverty level who are not otherwise covered by Medicaid or another program administered by a governmental entity, and to provide such services based on a sliding fee scale to all persons with incomes up to 200 percent of the federal poverty level who are not otherwise covered by Medicaid or another program administered by a governmental entity, except that eligibility may be limited to persons who
reside within a more limited area, as agreed to by the agency and the hospital.

(d) Contract with any federally qualified health center, if one exists within the agreed geopolitical boundaries, concerning the provision of primary care services, in order to guarantee delivery of services in a nonduplicative fashion, and to provide for referral arrangements, privileges, and admissions, as appropriate. The hospital shall agree to provide primary care services within 24 hours at an onsite or offsite facility to which all Medicaid recipients and persons eligible under this paragraph who do not require emergency room services are referred during normal daylight hours.

(e) Cooperate with the agency, the county, and other entities to ensure the provision of certain public health services, case management, referral and acceptance of patients, and sharing of epidemiological data, as the agency and the hospital find mutually necessary and desirable to promote and protect the public health within the agreed geopolitical boundaries.

(f) In cooperation with the county in which the hospital resides, develop a low-cost, outpatient, prepaid health care program to persons who are not eligible for the Medicaid program, and who reside within the area.

(g) Provide inpatient services to residents within the area who are not eligible for Medicaid or Medicare, and who do not have private health insurance, regardless of ability to pay, on the basis of available space, except that hospitals may not be prevented from establishing bill collection programs based on
ability to pay.

(h) Work with the Florida Healthy Kids Corporation, the Florida Health Care Purchasing Cooperative, and business health coalitions, as appropriate, to develop a feasibility study and plan to provide a low-cost comprehensive health insurance plan to persons who reside within the area and who do not have access to such a plan.

(i) Work with public health officials and other experts to provide community health education and prevention activities designed to promote healthy lifestyles and appropriate use of health services.

(j) Work with the local health council to develop a plan for promoting access to affordable health care services for all persons who reside within the area, including, but not limited to, public health services, primary care services, inpatient services, and affordable health insurance generally.

Any hospital that fails to comply with any of the provisions of this subsection, or any other contractual condition, may not receive payments under this section until full compliance is achieved.

Section 9. Paragraph (b) of subsection (16) and paragraph (a) of subsection (39) of section 409.912, Florida Statutes, are amended to read:

409.912 Cost-effective purchasing of health care.—The agency shall purchase goods and services for Medicaid recipients in the most cost-effective manner consistent with the delivery of quality medical care. To ensure that medical services are effectively utilized, the agency may, in any case, require a
confirmation or second physician’s opinion of the correct
diagnosis for purposes of authorizing future services under the
Medicaid program. This section does not restrict access to
emergency services or poststabilization care services as defined
in 42 C.F.R. part 438.114. Such confirmation or second opinion
shall be rendered in a manner approved by the agency. The agency
shall maximize the use of prepaid per capita and prepaid
aggregate fixed-sum basis services when appropriate and other
alternative service delivery and reimbursement methodologies,
including competitive bidding pursuant to s. 287.057, designed
to facilitate the cost-effective purchase of a case-managed
continuum of care. The agency shall also require providers to
minimize the exposure of recipients to the need for acute
inpatient, custodial, and other institutional care and the
inappropriate or unnecessary use of high-cost services. The
agency shall contract with a vendor to monitor and evaluate the
clinical practice patterns of providers in order to identify
trends that are outside the normal practice patterns of a
provider’s professional peers or the national guidelines of a
provider’s professional association. The vendor must be able to
provide information and counseling to a provider whose practice
patterns are outside the norms, in consultation with the agency,
to improve patient care and reduce inappropriate utilization.
The agency may mandate prior authorization, drug therapy
management, or disease management participation for certain
populations of Medicaid beneficiaries, certain drug classes, or
particular drugs to prevent fraud, abuse, overuse, and possible
dangerous drug interactions. The Pharmaceutical and Therapeutics
Committee shall make recommendations to the agency on drugs for

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which prior authorization is required. The agency shall inform
the Pharmaceutical and Therapeutics Committee of its decisions
regarding drugs subject to prior authorization. The agency is
authorized to limit the entities it contracts with or enrolls as
Medicaid providers by developing a provider network through
provider credentialing. The agency may competitively bid single-
source-provider contracts if procurement of goods or services
results in demonstrated cost savings to the state without
limiting access to care. The agency may limit its network based
on the assessment of beneficiary access to care, provider
availability, provider quality standards, time and distance
standards for access to care, the cultural competence of the
provider network, demographic characteristics of Medicaid
beneficiaries, practice and provider-to-beneficiary standards,
appointment wait times, beneficiary use of services, provider
turnover, provider profiling, provider licensure history,
previous program integrity investigations and findings, peer
review, provider Medicaid policy and billing compliance records,
icl临ical and medical record audits, and other factors. Providers
shall not be entitled to enrollment in the Medicaid provider
network. The agency shall determine instances in which allowing
Medicaid beneficiaries to purchase durable medical equipment and
other goods is less expensive to the Medicaid program than long-
term rental of the equipment or goods. The agency may establish
rules to facilitate purchases in lieu of long-term rentals in
order to protect against fraud and abuse in the Medicaid program
as defined in s. 409.913. The agency may seek federal waivers
necessary to administer these policies.

(16)
(b) The responsibility of the agency under this subsection includes the development of capabilities to identify actual and optimal practice patterns; patient and provider educational initiatives; methods for determining patient compliance with prescribed treatments; fraud, waste, and abuse prevention and detection programs; and beneficiary case management programs.

1. The practice pattern identification program shall evaluate practitioner prescribing patterns based on national and regional practice guidelines, comparing practitioners to their peer groups. The agency and its Drug Utilization Review Board shall consult with the Department of Health and a panel of practicing health care professionals consisting of the following: the Speaker of the House of Representatives and the President of the Senate shall each appoint three physicians licensed under chapter 458 or chapter 459; and the Governor shall appoint two pharmacists licensed under chapter 465 and one dentist licensed under chapter 466 who is an oral surgeon. Terms of the panel members shall expire at the discretion of the appointing official. The advisory panel shall be responsible for evaluating treatment guidelines and recommending ways to incorporate their use in the practice pattern identification program. Practitioners who are prescribing inappropriately or inefficiently, as determined by the agency, may have their prescribing of certain drugs subject to prior authorization or may be terminated from all participation in the Medicaid program.

2. The agency shall also develop educational interventions designed to promote the proper use of medications by providers
and beneficiaries.

3. The agency shall implement a pharmacy fraud, waste, and abuse initiative that may include a surety bond or letter of credit requirement for participating pharmacies, enhanced provider auditing practices, the use of additional fraud and abuse software, recipient management programs for beneficiaries inappropriately using their benefits, and other steps that will eliminate provider and recipient fraud, waste, and abuse. The initiative shall address enforcement efforts to reduce the number and use of counterfeit prescriptions.

4. By September 30, 2002, the agency may contract with an entity in the state to provide Medicaid providers with electronic access to Medicaid prescription refill data and information relating to the Medicaid Preferred Drug List. The agency may implement a wireless handheld clinical pharmacology drug information database for practitioners. The initiative shall be designed to enhance the agency’s efforts to reduce fraud, abuse, and errors in the prescription drug benefit program and to otherwise further the intent of this paragraph.

5. By April 1, 2006, the agency shall contract with an entity to design a database of clinical utilization information or electronic medical records for Medicaid providers. The database must be web-based and allow providers to review on a real-time basis the utilization of Medicaid services, including, but not limited to, physician office visits, inpatient and outpatient hospitalizations, laboratory and pathology services, radiological and other imaging services, dental care, and patterns of dispensing prescription drugs in order to coordinate care and identify potential fraud and abuse.
6. The agency may apply for any federal waivers needed to administer this paragraph.

   (39)(a) The agency shall implement a Medicaid prescribed-drug spending-control program that includes the following components:

   1. A Medicaid preferred drug list, which shall be a listing of cost-effective therapeutic options recommended by the Medicaid Pharmacy and Therapeutics Committee established pursuant to s. 409.91195 and adopted by the agency for each therapeutic class on the preferred drug list. At the discretion of the committee, and when feasible, the preferred drug list should include at least two products in a therapeutic class. The agency may post the preferred drug list and updates to the preferred drug list on an Internet website without following the rulemaking procedures of chapter 120. Antiretroviral agents are excluded from the preferred drug list. The agency shall also limit the amount of a prescribed drug dispensed to no more than a 34-day supply unless the drug products’ smallest marketed package is greater than a 34-day supply, or the drug is determined by the agency to be a maintenance drug in which case a 100-day maximum supply may be authorized. The agency may be authorized to seek any federal waivers necessary to implement these cost-control programs and to continue participation in the federal Medicaid rebate program, or alternatively to negotiate state-only manufacturer rebates. The agency may adopt rules to administer implement this subparagraph. The agency shall continue to provide unlimited contraceptive drugs and items. The agency must establish procedures to ensure that:

   a. There is a response to a request for prior consultation
by telephone or other telecommunication device within 24 hours
after receipt of a request for prior consultation; and

   b. A 72-hour supply of the drug prescribed is provided in
an emergency or when the agency does not provide a response
within 24 hours as required by sub-subparagraph a.

2. Reimbursement to pharmacies for Medicaid prescribed
drugs shall be set at the **lowest** lesser of: the average
wholesale price (AWP) minus 16.4 percent, the wholesaler
acquisition cost (WAC) plus 1.5 4.75 percent, the federal upper
limit (FUL), the state maximum allowable cost (SMAC), or the
usual and customary (UAC) charge billed by the provider.

3. The agency shall develop and implement a process for
managing the drug therapies of Medicaid recipients who are using
significant numbers of prescribed drugs each month. The
management process may include, but is not limited to,
comprehensive, physician-directed medical-record reviews, claims
analyses, and case evaluations to determine the medical
necessity and appropriateness of a patient’s treatment plan and
drug therapies. The agency may contract with a private
organization to provide drug-program-management services. The
Medicaid drug benefit management program shall include
initiatives to manage drug therapies for HIV/AIDS patients,
patients using 20 or more unique prescriptions in a 180-day
period, and the top 1,000 patients in annual spending. The
agency shall enroll any Medicaid recipient in the drug benefit
management program if he or she meets the specifications of this
provision and is not enrolled in a Medicaid health maintenance
organization.

4. The agency may limit the size of its pharmacy network
based on need, competitive bidding, price negotiations, credentialing, or similar criteria. The agency shall give special consideration to rural areas in determining the size and location of pharmacies included in the Medicaid pharmacy network. A pharmacy credentialing process may include criteria such as a pharmacy’s full-service status, location, size, patient educational programs, patient consultation, disease management services, and other characteristics. The agency may impose a moratorium on Medicaid pharmacy enrollment if when it is determined that it has a sufficient number of Medicaid-participating providers. The agency must allow dispensing practitioners to participate as a part of the Medicaid pharmacy network regardless of the practitioner’s proximity to any other entity that is dispensing prescription drugs under the Medicaid program. A dispensing practitioner must meet all credentialing requirements applicable to his or her practice, as determined by the agency.

5. The agency shall develop and implement a program that requires Medicaid practitioners who prescribe drugs to use a counterfeit-proof prescription pad for Medicaid prescriptions. The agency shall require the use of standardized counterfeit-proof prescription pads by Medicaid-participating prescribers or prescribers who write prescriptions for Medicaid recipients. The agency may implement the program in targeted geographic areas or statewide.

6. The agency may enter into arrangements that require manufacturers of generic drugs prescribed to Medicaid recipients to provide rebates of at least 15.1 percent of the average manufacturer price for the manufacturer’s generic products.
These arrangements shall require that if a generic-drug manufacturer pays federal rebates for Medicaid-reimbursed drugs at a level below 15.1 percent, the manufacturer must provide a supplemental rebate to the state in an amount necessary to achieve a 15.1-percent rebate level.

7. The agency may establish a preferred drug list as described in this subsection, and, pursuant to the establishment of such preferred drug list, it is authorized to negotiate supplemental rebates from manufacturers that are in addition to those required by Title XIX of the Social Security Act and at no less than 14 percent of the average manufacturer price as defined in 42 U.S.C. s. 1936 on the last day of a quarter unless the federal or supplemental rebate, or both, equals or exceeds 29 percent. There is no upper limit on the supplemental rebates the agency may negotiate. The agency may determine that specific products, brand-name or generic, are competitive at lower rebate percentages. Agreement to pay the minimum supplemental rebate percentage will guarantee a manufacturer that the Medicaid Pharmaceutical and Therapeutics Committee will consider a product for inclusion on the preferred drug list. However, a pharmaceutical manufacturer is not guaranteed placement on the preferred drug list by simply paying the minimum supplemental rebate. Agency decisions will be made on the clinical efficacy of a drug and recommendations of the Medicaid Pharmaceutical and Therapeutics Committee, as well as the price of competing products minus federal and state rebates. The agency may be authorized to contract with an outside agency or contractor to conduct negotiations for supplemental rebates. For the purposes of this section, the term “supplemental rebates” means cash.
rebates. Effective July 1, 2004, Value-added programs as a substitution for supplemental rebates are prohibited. The agency may be authorized to seek any federal waivers to implement this initiative.

8. The agency for Health Care Administration shall expand home delivery of pharmacy products. To assist Medicaid recipients in securing their prescriptions and reduce program costs, the agency shall expand its current mail-order-pharmacy diabetes-supply program to include all generic and brand-name drugs used by Medicaid recipients with diabetes. Medicaid recipients in the current program may obtain nondiabetes drugs on a voluntary basis. This initiative is limited to the geographic area covered by the current contract. The agency may seek and implement any federal waivers necessary to implement this subparagraph.

9. The agency shall limit to one dose per month any drug prescribed to treat erectile dysfunction.

10.a. The agency may implement a Medicaid behavioral drug management system. The agency may contract with a vendor that has experience in operating behavioral drug management systems to implement this program. The agency may be authorized to seek federal waivers to implement this program.

b. The agency, in conjunction with the Department of Children and Family Services, may implement the Medicaid behavioral drug management system that is designed to improve the quality of care and behavioral health prescribing practices based on best practice guidelines, improve patient adherence to medication plans, reduce clinical risk, and lower prescribed drug costs and the rate of inappropriate spending on Medicaid
behavioral drugs. The program may include the following elements:

(I) Provide for the development and adoption of best practice guidelines for behavioral health-related drugs such as antipsychotics, antidepressants, and medications for treating bipolar disorders and other behavioral conditions; translate them into practice; review behavioral health prescribers and compare their prescribing patterns to a number of indicators that are based on national standards; and determine deviations from best practice guidelines.

(II) Implement processes for providing feedback to and educating prescribers using best practice educational materials and peer-to-peer consultation.

(III) Assess Medicaid beneficiaries who are outliers in their use of behavioral health drugs with regard to the numbers and types of drugs taken, drug dosages, combination drug therapies, and other indicators of improper use of behavioral health drugs.

(IV) Alert prescribers to patients who fail to refill prescriptions in a timely fashion, are prescribed multiple same-class behavioral health drugs, and may have other potential medication problems.

(V) Track spending trends for behavioral health drugs and deviation from best practice guidelines.

(VI) Use educational and technological approaches to promote best practices, educate consumers, and train prescribers in the use of practice guidelines.

(VII) Disseminate electronic and published materials.

(VIII) Hold statewide and regional conferences.
(IX) Implement a disease management program with a model quality-based medication component for severely mentally ill individuals and emotionally disturbed children who are high users of care.

11. The agency shall implement a Medicaid prescription drug management system.

a. The agency may contract with a vendor that has experience in operating prescription drug management systems in order to implement this system. Any management system that is implemented in accordance with this subparagraph must rely on cooperation between physicians and pharmacists to determine appropriate practice patterns and clinical guidelines to improve the prescribing, dispensing, and use of drugs in the Medicaid program. The agency may seek federal waivers to implement this program.

b. The drug management system must be designed to improve the quality of care and prescribing practices based on best practice guidelines, improve patient adherence to medication plans, reduce clinical risk, and lower prescribed drug costs and the rate of inappropriate spending on Medicaid prescription drugs. The program must:

(I) Provide for the development and adoption of best practice guidelines for the prescribing and use of drugs in the Medicaid program, including translating best practice guidelines into practice; reviewing prescriber patterns and comparing them to indicators that are based on national standards and practice patterns of clinical peers in their community, statewide, and nationally; and determine deviations from best practice guidelines.
(II) Implement processes for providing feedback to and educating prescribers using best practice educational materials and peer-to-peer consultation.

(III) Assess Medicaid recipients who are outliers in their use of a single or multiple prescription drugs with regard to the numbers and types of drugs taken, drug dosages, combination drug therapies, and other indicators of improper use of prescription drugs.

(IV) Alert prescribers to recipients patients who fail to refill prescriptions in a timely fashion, are prescribed multiple drugs that may be redundant or contraindicated, or may have other potential medication problems.

(V) Track spending trends for prescription drugs and deviation from best practice guidelines.

(VI) Use educational and technological approaches to promote best practices, educate consumers, and train prescribers in the use of practice guidelines.

(VII) Disseminate electronic and published materials.

(VIII) Hold statewide and regional conferences.

(IX) Implement disease management programs in cooperation with physicians and pharmacists, along with a model quality-based medication component for individuals having chronic medical conditions.

12. The agency may be authorized to contract for drug rebate administration, including, but not limited to, calculating rebate amounts, invoicing manufacturers, negotiating disputes with manufacturers, and maintaining a database of rebate collections.

13. The agency may specify the preferred daily dosing form
or strength for the purpose of promoting best practices with regard to the prescribing of certain drugs as specified in the General Appropriations Act and ensuring cost-effective prescribing practices.

14. The agency may require prior authorization for Medicaid-covered prescribed drugs. The agency may, but is not required to, prior authorize the use of a product:
   a. For an indication not approved in labeling;
   b. To comply with certain clinical guidelines; or
   c. If the product has the potential for overuse, misuse, or abuse.

The agency may require the prescribing professional to provide information about the rationale and supporting medical evidence for the use of a drug. The agency may post prior authorization criteria and protocol and updates to the list of drugs that are subject to prior authorization on an Internet website without amending its rule or engaging in additional rulemaking.

15. The agency, in conjunction with the Pharmaceutical and Therapeutics Committee, may require age-related prior authorizations for certain prescribed drugs. The agency may preauthorize the use of a drug for a recipient who may not meet the age requirement or may exceed the length of therapy for use of this product as recommended by the manufacturer and approved by the Food and Drug Administration. Prior authorization may require the prescribing professional to provide information about the rationale and supporting medical evidence for the use of a drug.

16. The agency shall implement a step-therapy prior
authorization approval process for medications excluded from the preferred drug list. Medications listed on the preferred drug list must be used within the previous 12 months before prior to the alternative medications that are not listed. The step-therapy prior authorization may require the prescriber to use the medications of a similar drug class or for a similar medical indication unless contraindicated in the Food and Drug Administration labeling. The trial period between the specified steps may vary according to the medical indication. The step-therapy approval process shall be developed in accordance with the committee as stated in s. 409.91195(7) and (8). A drug product may be approved without meeting the step-therapy prior authorization criteria if the prescribing physician provides the agency with additional written medical or clinical documentation that the product is medically necessary because:

a. There is not a drug on the preferred drug list to treat the disease or medical condition which is an acceptable clinical alternative;

b. The alternatives have been ineffective in the treatment of the beneficiary’s disease; or

c. Based on historic evidence and known characteristics of the patient and the drug, the drug is likely to be ineffective, or the number of doses have been ineffective.

The agency shall work with the physician to determine the best alternative for the patient. The agency may adopt rules waiving the requirements for written clinical documentation for specific drugs in limited clinical situations.

17. The agency shall implement a return and reuse program
for drugs dispensed by pharmacies to institutional recipients, which includes payment of a $5 restocking fee for the implementation and operation of the program. The return and reuse program shall be implemented electronically and in a manner that promotes efficiency. The program must permit a pharmacy to exclude drugs from the program if it is not practical or cost-effective for the drug to be included and must provide for the return to inventory of drugs that cannot be credited or returned in a cost-effective manner. The agency shall determine if the program has reduced the amount of Medicaid prescription drugs which are destroyed on an annual basis and if there are additional ways to ensure more prescription drugs are not destroyed which could safely be reused. The agency’s conclusion and recommendations shall be reported to the Legislature by December 1, 2005.

Section 10. Paragraph (a) of subsection (2) of section 409.9122, Florida Statutes, is amended to read:

409.9122 Mandatory Medicaid managed care enrollment; programs and procedures.—

(2)(a) The agency shall enroll all Medicaid recipients in a managed care plan or MediPass all Medicaid recipients, except those Medicaid recipients who are: in an institution, receiving a Medicaid nonpoverty medical subsidy, enrolled in the Medicaid medically needy Program, or eligible for both Medicaid and Medicare. Upon enrollment, recipients may individuals will be able to change their managed care option during the 90-day opt out period required by federal Medicaid regulations. The agency may is authorized to seek the necessary Medicaid state plan amendment to implement this policy. However, to the extent
1. If permitted by federal law, the agency may enroll in a managed care plan or MediPass a Medicaid recipient who is exempt from mandatory managed care enrollment in a managed care plan or MediPass if, provided that:
   a. The recipient’s decision to enroll in a managed care plan or MediPass is voluntary;
   b. If the recipient chooses to enroll in a managed care plan, the agency has determined that the managed care plan provides specific programs and services that address the special health needs of the recipient; and
   c. The agency receives any necessary waivers from the federal Centers for Medicare and Medicaid Services.

2. The agency shall develop rules to establish policies by which exceptions to the mandatory managed care enrollment requirement may be made on a case-by-case basis. The rules shall include the specific criteria to be applied when determining whether to exempt a recipient from mandatory enrollment in a managed care plan or MediPass.

3. School districts participating in the certified school match program pursuant to ss. 409.908(21) and 1011.70 shall be reimbursed by Medicaid, subject to the limitations of s. 1011.70(1), for a Medicaid-eligible child participating in the services authorized in s. 1011.70, as provided for in s. 409.9071, regardless of whether the child is enrolled in MediPass or a managed care plan. Managed care plans shall make a good faith effort to execute agreements with school districts regarding the coordinated provision of services authorized under s. 1011.70.
4. County health departments delivering school-based services pursuant to ss. 381.0056 and 381.0057 shall be reimbursed by Medicaid for the federal share for a Medicaid-eligible child who receives Medicaid-covered services in a school setting, regardless of whether the child is enrolled in MediPass or a managed care plan. Managed care plans shall make a good faith effort to execute agreements with county health departments that coordinate the provision of services to a Medicaid-eligible child. To ensure continuity of care for Medicaid patients, the agency, the Department of Health, and the Department of Education shall develop procedures for ensuring that a student’s managed care plan or MediPass provider receives information relating to services provided in accordance with ss. 381.0056, 381.0057, 409.9071, and 1011.70.

Section 11. Paragraph (a) of subsection (1) of section 409.915, Florida Statutes, is amended to read:

409.915 County contributions to Medicaid.—Although the state is responsible for the full portion of the state share of the matching funds required for the Medicaid program, in order to acquire a certain portion of these funds, the state shall charge the counties for certain items of care and service as provided in this section.

(1) Each county shall participate in the following items of care and service:

(a) For both health maintenance members and fee-for-service beneficiaries, payments for inpatient hospitalization in excess of 10 days, but not in excess of 45 days, with the exception of pregnant women and children whose income is greater than in
excess of the federal poverty level and who do not receive a
Medicaid nonpoverty medical subsidy under s. 409.904(2)
participate in the Medicaid medically needy Program, and for
adult lung transplant services.
Section 12. Subsections (1) and (2) of section 409.9301,
Florida Statutes, are amended to read:
409.9301 Pharmaceutical expense assistance.—
(1) PROGRAM ESTABLISHED.—A program is established in the
agency for Health Care Administration to provide pharmaceutical
expense assistance to individuals diagnosed with cancer or
individuals who have obtained received organ transplants who
received a Medicaid nonpoverty medical subsidy before were
medically needy recipients prior to January 1, 2006.
(2) ELIGIBILITY.—Eligibility for the program is limited to
an individual who:
(a) Is a resident of this state;
(b) Was a Medicaid recipient who received a Medicaid
nonpoverty medical subsidy before under the Florida Medicaid
medically needy program prior to January 1, 2006;
(c) Is eligible for Medicare;
(d) Is a cancer patient or an organ transplant recipient;
and
(e) Requests to be enrolled in the program.
Section 13. This act shall take effect June 30, 2011.