Bill No. CS for SB 2034, 1st Eng.

Amendment No. ____

	CHAMBER ACTION <u>Senate</u> <u>House</u>
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11	Senator Silver moved the following amendment:
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13	Senate Amendment (with title amendment)
14	On page 66, between lines 22 and 23,
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16	insert:
17	Section 38. Subsection (37) is added to section
18	409.912, Florida Statutes, to read:
19	409.912 Cost-effective purchasing of health careThe
20	agency shall purchase goods and services for Medicaid
21	recipients in the most cost-effective manner consistent with
22	the delivery of quality medical care. The agency shall
23	maximize the use of prepaid per capita and prepaid aggregate
24	fixed-sum basis services when appropriate and other
25	alternative service delivery and reimbursement methodologies,
26	including competitive bidding pursuant to s. 287.057, designed
27	to facilitate the cost-effective purchase of a case-managed
28	continuum of care. The agency shall also require providers to
29	minimize the exposure of recipients to the need for acute
30	inpatient, custodial, and other institutional care and the
31	inappropriate or unnecessary use of high-cost services.

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(37)(a) The agency shall implement a Medicaid prescribed-drug spending-control program that includes the following components: 1. Medicaid prescribed-drug coverage for brand-name drugs for adult Medicaid recipients not residing in nursing homes or other institutions is limited to the dispensing of four brand-name drugs per month per recipient. Children and institutionalized adults are exempt from this restriction. Antiretroviral agents are excluded from this limitation. No requirements for prior authorization or other restrictions on medications used to treat mental illnesses such as schizophrenia, severe depression, or bipolar disorder may be imposed on Medicaid recipients. Medications that will be available without restriction for persons with mental illnesses include atypical antipsychotic medications, conventional antipsychotic medications, selective serotonin re-uptake inhibitors, and other medications used for the treatment of serious mental illnesses. The agency shall also limit the amount of a prescribed drug dispensed to no more than a 34-day supply. The agency shall continue to provide unlimited generic drugs, contraceptive drugs and items, and diabetic supplies. The agency may authorize exceptions to the brand-name-drug restriction, based upon the treatment needs of the patients, only when such exceptions are based on prior

a. There will be 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

consultation provided by the agency or an agency contractor,

but the agency must establish procedures to ensure that:

b. A 72-hour supply of the drug prescribed will be

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 average wholesale price less 13.25 percent.
- 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.
- 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 when it is determined that it has a sufficient number of Medicaid-participating providers.
- 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. 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. If a generic-drug manufacturer raises its price in excess of the Consumer Price Index (Urban), the excess amount shall be included in the supplemental rebate to the state.
- (b) The agency shall implement this subsection to the extent that funds are appropriated to administer the Medicaid prescribed-drug spending-control program. The agency may contract all or any part of this program to private organizations.
- (c) The agency shall submit a report to the Governor, the President of the Senate, and the Speaker of the House of Representatives by January 15 of each year. The report must include, but need not be limited to, the progress made in implementing Medicaid cost-containment measures and their effect on Medicaid prescribed-drug expenditures.
- Section 39. There is created a Medicaid Pharmaceutical and Therapeutics Committee. The committee shall develop and implement a voluntary Medicaid preferred prescribed drug designation program. The program shall provide information to Medicaid providers on medically appropriate and cost efficient prescription drug therapies through the development and

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publication of a voluntary Medicaid preferred prescribed-drug
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    list.
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          (1) The Medicaid Pharmaceutical and Therapeutics
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    Committee shall be comprised of nine members appointed as
    follows: one practicing physician licensed under chapter 458,
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    Florida Statutes, appointed by the Speaker of the House of
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    Representatives from a list of recommendations from the
   Florida Medical Association; one practicing physician licensed
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   under chapter 459, Florida Statutes, appointed by the Speaker
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    of the House of Representatives from a list of recommendations
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    from the Florida Osteopathic Medical Association; one
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   practicing physician licensed under chapter 458, Florida
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    Statutes, appointed by the President of the Senate from a list
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    of recommendations from the Florida Academy of Family
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    Physicians; one practicing podiatric physician licensed under
    chapter 461, Florida Statutes, appointed by the President of
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    the Florida Senate from a list of recommendations from the
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    Florida Podiatric Medical Association; one trauma surgeon
    licensed under chapter 458, Florida Statutes, appointed by the
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    Speaker of the House of Representatives from a list of
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    recommendations from the American College of Surgeons; one
   practicing dentist licensed under chapter 466, Florida
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    Statutes, appointed by the President of the Senate from a list
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    of recommendations from the Florida Dental Association; one
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    practicing pharmacist licensed under chapter 465, Florida
    Statutes, appointed by the Governor from a list of
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    recommendations from the Florida Pharmacy Association; one
   practicing pharmacist licensed under chapter 465, Florida
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    Statutes, appointed by the Governor from a list of
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    recommendations from the Florida Society of Health System
31 | Pharmacists; and one health care professional with expertise
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in clinical pharmacology appointed by the Governor from a list
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    of recommendations from the Pharmaceutical Research and
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    Manufacturers Association. The members shall be appointed to
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    serve for terms of 2 years from the date of their appointment.
    Members may be appointed to more than one term. The Agency for
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    Health Care Administration shall serve as staff for the
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    committee and assist them with all ministerial duties.
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          (2) Upon recommendation by the committee, the Agency
    for Health Care Administration shall establish the voluntary
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    Medicaid preferred prescribed-drug list. Upon further
    recommendation by the committee, the agency shall add to,
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    delete from, or modify the list. The committee shall also
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    review requests for additions to, deletions from, or
   modifications of the list. The list shall be adopted by the
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   committee in consultation with medical specialists, when
    appropriate, using the following criteria: use of the list
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    shall be voluntary by providers and the list must provide for
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   medically appropriate drug therapies for Medicaid patients
    which achieve cost savings in the Medicaid program.
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          (3) The Agency for Health Care Administration shall
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    publish and disseminate the voluntary Medicaid preferred
   prescribed drug list to all Medicaid providers in the state.
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    (Redesignate subsequent sections.)
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    ======= T I T L E A M E N D M E N T =========
28
    And the title is amended as follows:
           On page 6, line 4, following the semicolon
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30
31 insert:
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amending s. 409.912, F.S., relating to cost-effective purchasing of health care under the Medicaid program; requiring the agency to implement a Medicaid prescribed-drug spending-control program; specifying program components; providing for implementation to the extent funds are appropriated; authorizing contracts; requiring an annual report; creating the Medicaid Pharmaceutical Therapeutics Committee; providing for membership; providing for the adoption of a voluntary preferred prescribed-drug list by the committee;