

By Senator Harrell

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A bill to be entitled
An act relating to step-therapy protocols; amending s.
409.901, F.S.; defining the term "serious mental
illness"; amending s. 409.912, F.S.; requiring the
Agency for Health Care Administration to approve drug
products for Medicaid recipients for the treatment of
serious mental illness without step-therapy prior
authorization under certain circumstances; amending s.
409.910, F.S.; conforming a cross-reference; directing
the agency to include rate impacts resulting from the
act in certain rates that become effective on a
specified date; providing effective dates.

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

Section 1. Present subsections (27) and (28) of section
409.901, Florida Statutes, are redesignated as subsections (28)
and (29), respectively, and a new subsection (27) is added to
that section, to read:

409.901 Definitions; ss. 409.901-409.920.—As used in ss.
409.901-409.920, except as otherwise specifically provided, the
term:

(27) "Serious mental illness" means any of the following
psychiatric disorders as defined in the Diagnostic and
Statistical Manual of Mental Disorders, Fifth Edition, published
by the American Psychiatric Association:

(a) Bipolar disorders, including hypomanic, manic,
depressive, and mixed-feature episodes.

(b) Depression in childhood or adolescence.

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30 (c) Major depressive disorders, including single and
31 recurrent depressive episodes.

32 (d) Obsessive-compulsive disorders.

33 (e) Paranoid personality disorder or other psychotic
34 disorders.

35 (f) Schizoaffective disorders, including bipolar or
36 depressive symptoms.

37 (g) Schizophrenia.

38 Section 2. Paragraph (a) of subsection (5) of section
39 409.912, Florida Statutes, is amended to read:

40 409.912 Cost-effective purchasing of health care.—The
41 agency shall purchase goods and services for Medicaid recipients
42 in the most cost-effective manner consistent with the delivery
43 of quality medical care. To ensure that medical services are
44 effectively utilized, the agency may, in any case, require a
45 confirmation or second physician's opinion of the correct
46 diagnosis for purposes of authorizing future services under the
47 Medicaid program. This section does not restrict access to
48 emergency services or poststabilization care services as defined
49 in 42 C.F.R. s. 438.114. Such confirmation or second opinion
50 shall be rendered in a manner approved by the agency. The agency
51 shall maximize the use of prepaid per capita and prepaid
52 aggregate fixed-sum basis services when appropriate and other
53 alternative service delivery and reimbursement methodologies,
54 including competitive bidding pursuant to s. 287.057, designed
55 to facilitate the cost-effective purchase of a case-managed
56 continuum of care. The agency shall also require providers to
57 minimize the exposure of recipients to the need for acute
58 inpatient, custodial, and other institutional care and the

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59 inappropriate or unnecessary use of high-cost services. The
60 agency shall contract with a vendor to monitor and evaluate the
61 clinical practice patterns of providers in order to identify
62 trends that are outside the normal practice patterns of a
63 provider's professional peers or the national guidelines of a
64 provider's professional association. The vendor must be able to
65 provide information and counseling to a provider whose practice
66 patterns are outside the norms, in consultation with the agency,
67 to improve patient care and reduce inappropriate utilization.
68 The agency may mandate prior authorization, drug therapy
69 management, or disease management participation for certain
70 populations of Medicaid beneficiaries, certain drug classes, or
71 particular drugs to prevent fraud, abuse, overuse, and possible
72 dangerous drug interactions. The Pharmaceutical and Therapeutics
73 Committee shall make recommendations to the agency on drugs for
74 which prior authorization is required. The agency shall inform
75 the Pharmaceutical and Therapeutics Committee of its decisions
76 regarding drugs subject to prior authorization. The agency is
77 authorized to limit the entities it contracts with or enrolls as
78 Medicaid providers by developing a provider network through
79 provider credentialing. The agency may competitively bid single-
80 source-provider contracts if procurement of goods or services
81 results in demonstrated cost savings to the state without
82 limiting access to care. The agency may limit its network based
83 on the assessment of beneficiary access to care, provider
84 availability, provider quality standards, time and distance
85 standards for access to care, the cultural competence of the
86 provider network, demographic characteristics of Medicaid
87 beneficiaries, practice and provider-to-beneficiary standards,

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88 appointment wait times, beneficiary use of services, provider
89 turnover, provider profiling, provider licensure history,
90 previous program integrity investigations and findings, peer
91 review, provider Medicaid policy and billing compliance records,
92 clinical and medical record audits, and other factors. Providers
93 are not entitled to enrollment in the Medicaid provider network.
94 The agency shall determine instances in which allowing Medicaid
95 beneficiaries to purchase durable medical equipment and other
96 goods is less expensive to the Medicaid program than long-term
97 rental of the equipment or goods. The agency may establish rules
98 to facilitate purchases in lieu of long-term rentals in order to
99 protect against fraud and abuse in the Medicaid program as
100 defined in s. 409.913. The agency may seek federal waivers
101 necessary to administer these policies.

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

105 1. A Medicaid preferred drug list, which shall be a listing
106 of cost-effective therapeutic options recommended by the
107 Medicaid Pharmacy and Therapeutics Committee established
108 pursuant to s. 409.91195 and adopted by the agency for each
109 therapeutic class on the preferred drug list. At the discretion
110 of the committee, and when feasible, the preferred drug list
111 should include at least two products in a therapeutic class. The
112 agency may post the preferred drug list and updates to the list
113 on an Internet website without following the rulemaking
114 procedures of chapter 120. Antiretroviral agents are excluded
115 from the preferred drug list. The agency shall also limit the
116 amount of a prescribed drug dispensed to no more than a 34-day

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117 supply unless the drug products' smallest marketed package is
118 greater than a 34-day supply, or the drug is determined by the
119 agency to be a maintenance drug, in which case a 100-day maximum
120 supply may be authorized. The agency may seek any federal
121 waivers necessary to implement these cost-control programs and
122 to continue participation in the federal Medicaid rebate
123 program, or alternatively to negotiate state-only manufacturer
124 rebates. The agency may adopt rules to administer this
125 subparagraph. The agency shall continue to provide unlimited
126 contraceptive drugs and items. The agency must establish
127 procedures to ensure that:

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

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

134 2. A provider of prescribed drugs is reimbursed in an
135 amount not to exceed the lesser of the actual acquisition cost
136 based on the Centers for Medicare and Medicaid Services National
137 Average Drug Acquisition Cost pricing files plus a professional
138 dispensing fee, the wholesale acquisition cost plus a
139 professional dispensing fee, the state maximum allowable cost
140 plus a professional dispensing fee, or the usual and customary
141 charge billed by the provider.

142 3. The agency shall develop and implement a process for
143 managing the drug therapies of Medicaid recipients who are using
144 significant numbers of prescribed drugs each month. The
145 management process may include, but is not limited to,

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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 must ~~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 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

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175 requirements applicable to his or her practice, as determined by
176 the agency.

177 5. The agency shall develop and implement a program that
178 requires Medicaid practitioners who issue written prescriptions
179 for medicinal drugs to use a counterfeit-proof prescription pad
180 for Medicaid prescriptions. The agency shall require the use of
181 standardized counterfeit-proof prescription pads by prescribers
182 who issue written prescriptions for Medicaid recipients. The
183 agency may implement the program in targeted geographic areas or
184 statewide.

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

194 7. The agency may establish a preferred drug list as
195 described in this subsection, and, pursuant to the establishment
196 of such preferred drug list, negotiate supplemental rebates from
197 manufacturers that are in addition to those required by Title
198 XIX of the Social Security Act and at no less than 14 percent of
199 the average manufacturer price as defined in 42 U.S.C. s. 1936
200 on the last day of a quarter unless the federal or supplemental
201 rebate, or both, equals or exceeds 29 percent. There is no upper
202 limit on the supplemental rebates the agency may negotiate. The
203 agency may determine that specific products, brand-name or

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generic, are competitive at lower rebate percentages. Agreement to pay the minimum supplemental rebate percentage guarantees 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 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. Value-added programs as a substitution for supplemental rebates are prohibited. The agency may seek any federal waivers to implement this initiative.

8.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 seek federal waivers to implement this program.

b. The agency, in conjunction with the Department of Children and Families, 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

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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

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quality-based medication component for severely mentally ill individuals and emotionally disturbed children who are high users of care.

9. 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 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

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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 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.

10. The agency may 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.

11. 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.

12. The agency may require prior authorization for Medicaid-covered prescribed drugs. The agency may 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

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for the use of a drug. The agency shall post prior authorization, step-edit criteria and protocol, and updates to the list of drugs that are subject to prior authorization on the agency's Internet website within 21 days after the prior authorization and step-edit criteria and protocol and updates are approved by the agency. For purposes of this subparagraph, the term "step-edit" means an automatic electronic review of certain medications subject to prior authorization.

13. 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.

14. 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 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 must ~~shall~~ be developed in accordance

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with the committee as stated in s. 409.91195(7) and (8). A drug product may be approved, or, in the case of a drug product for the treatment of a serious mental illness, must 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;

c. The drug product or medication of a similar drug class is prescribed for the treatment of a serious mental illness ~~schizophrenia or schizotypal or delusional disorders~~; prior authorization has been granted previously for the prescribed drug; and the medication was dispensed to the patient during the previous 12 months; or

d. Based on historical 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.

15. 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

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implementation and operation of the program. The return and reuse program must ~~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 whether ~~if~~ the program has reduced the amount of Medicaid prescription drugs which are destroyed on an annual basis and whether ~~if~~ there are additional ways to ensure more prescription drugs are not destroyed and ~~which~~ could safely be reused.

Section 3. Paragraph (a) of subsection (20) of section 409.910, Florida Statutes, is amended to read:

409.910 Responsibility for payments on behalf of Medicaid-eligible persons when other parties are liable.—

(20) (a) Entities providing health insurance as defined in s. 624.603, health maintenance organizations and prepaid health clinics as defined in chapter 641, and, on behalf of their clients, third-party administrators, pharmacy benefits managers, and any other third parties, as defined in s. 409.901 ~~s. 409.901(27)~~, which are legally responsible for payment of a claim for a health care item or service as a condition of doing business in this ~~the~~ state or providing coverage to residents of this state, shall provide such records and information as are necessary to accomplish the purpose of this section, unless such requirement results in an unreasonable burden.

Section 4. The Agency for Health Care Administration is directed to include the rate impact of this act in the Medicaid

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407 managed medical assistance program and long-term care managed
408 care program rates that become effective on October 1, 2026.
409 This section shall take effect upon this act becoming a law.

410 Section 5. Except as otherwise expressly provided in this
411 act and except for this section, which shall take effect upon
412 this act becoming a law, this act shall take effect October 1,
413 2026.