

**SENATE  
STATE OF MINNESOTA  
NINETY-THIRD SESSION**

**S.F. No. 4368**

(SENATE AUTHORS: HOFFMAN)

DATE  
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D-PG

Introduction and first reading  
Referred to Health and Human Services

OFFICIAL STATUS

1.1 A bill for an act  
1.2 relating to human services; expanding prior authorization mental health carve out;  
1.3 amending Minnesota Statutes 2023 Supplement, section 256B.0625, subdivision  
1.4 13f.

1.5 BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF MINNESOTA:

1.6 Section 1. Minnesota Statutes 2023 Supplement, section 256B.0625, subdivision 13f, is  
1.7 amended to read:

1.8 Subd. 13f. **Prior authorization.** (a) The Formulary Committee shall review and  
1.9 recommend drugs which require prior authorization. The Formulary Committee shall  
1.10 establish general criteria to be used for the prior authorization of brand-name drugs for  
1.11 which generically equivalent drugs are available, but the committee is not required to review  
1.12 each brand-name drug for which a generically equivalent drug is available.

1.13 (b) Prior authorization may be required by the commissioner before certain formulary  
1.14 drugs are eligible for payment. The Formulary Committee may recommend drugs for prior  
1.15 authorization directly to the commissioner. The commissioner may also request that the  
1.16 Formulary Committee review a drug for prior authorization. Before the commissioner may  
1.17 require prior authorization for a drug:

1.18 (1) the commissioner must provide information to the Formulary Committee on the  
1.19 impact that placing the drug on prior authorization may have on the quality of patient care  
1.20 and on program costs, information regarding whether the drug is subject to clinical abuse  
1.21 or misuse, and relevant data from the state Medicaid program if such data is available;

1.22 (2) the Formulary Committee must review the drug, taking into account medical and  
1.23 clinical data and the information provided by the commissioner; and

2.1 (3) the Formulary Committee must hold a public forum and receive public comment for  
2.2 an additional 15 days.

2.3 The commissioner must provide a 15-day notice period before implementing the prior  
2.4 authorization.

2.5 (c) Except as provided in subdivision 13j, prior authorization or other utilization  
2.6 management controls shall not be required or utilized for:

2.7 (1) any atypical antipsychotic drug prescribed for the treatment of mental illness ~~if;~~ or

2.8 (2) any Federal Drug Administration-approved medicine defined in the most recent  
2.9 edition of the Diagnostic and Statistical Manual of Mental Disorders published by the  
2.10 American Psychiatric Association that is indicated for the treatment of a mental disorder  
2.11 or condition that results in a serious functional impairment that substantially interferes with  
2.12 or limits one or more major life activities.

2.13 ~~(1) there is no generically equivalent drug available; and~~

2.14 ~~(2) the drug was initially prescribed for the recipient prior to July 1, 2003; or~~

2.15 ~~(3) the drug is part of the recipient's current course of treatment.~~

2.16 This paragraph applies to any multistate preferred drug list or supplemental drug rebate  
2.17 program established or administered by the commissioner. Prior authorization shall  
2.18 automatically be granted for 60 days for brand name drugs prescribed for treatment of mental  
2.19 illness within 60 days of when a generically equivalent drug becomes available, provided  
2.20 that the brand name drug was part of the recipient's course of treatment at the time the  
2.21 generically equivalent drug became available.

2.22 For purposes of this paragraph, "utilization management controls" means a set of formal  
2.23 techniques used by a health carrier or prescription drug utilization management entity that  
2.24 are designed to monitor the use of prescription drugs or evaluate their medical necessity,  
2.25 appropriateness, efficacy, and efficiency, including but not limited to prior authorization  
2.26 processes and step therapy protocols as defined in section 62Q.1841.

2.27 (d) Prior authorization must not be required for liquid methadone if only one version of  
2.28 liquid methadone is available. If more than one version of liquid methadone is available,  
2.29 the commissioner shall ensure that at least one version of liquid methadone is available  
2.30 without prior authorization.

2.31 (e) Prior authorization may be required for an oral liquid form of a drug, except as  
2.32 described in paragraph (d). A prior authorization request under this paragraph must be

3.1 automatically approved within 24 hours if the drug is being prescribed for a Food and Drug  
3.2 Administration-approved condition for a patient who utilizes an enteral tube for feedings  
3.3 or medication administration, even if the patient has current or prior claims for pills for that  
3.4 condition. If more than one version of the oral liquid form of a drug is available, the  
3.5 commissioner may select the version that is able to be approved for a Food and Drug  
3.6 Administration-approved condition for a patient who utilizes an enteral tube for feedings  
3.7 or medication administration. This paragraph applies to any multistate preferred drug list  
3.8 or supplemental drug rebate program established or administered by the commissioner. The  
3.9 commissioner shall design and implement a streamlined prior authorization form for patients  
3.10 who utilize an enteral tube for feedings or medication administration and are prescribed an  
3.11 oral liquid form of a drug. The commissioner may require prior authorization for brand  
3.12 name drugs whenever a generically equivalent product is available, even if the prescriber  
3.13 specifically indicates "dispense as written-brand necessary" on the prescription as required  
3.14 by section 151.21, subdivision 2.

3.15 (f) Notwithstanding this subdivision, the commissioner may automatically require prior  
3.16 authorization, for a period not to exceed 180 days, for any drug that is approved by the  
3.17 United States Food and Drug Administration on or after July 1, 2005. The 180-day period  
3.18 begins no later than the first day that a drug is available for shipment to pharmacies within  
3.19 the state. The Formulary Committee shall recommend to the commissioner general criteria  
3.20 to be used for the prior authorization of the drugs, but the committee is not required to  
3.21 review each individual drug. In order to continue prior authorizations for a drug after the  
3.22 180-day period has expired, the commissioner must follow the provisions of this subdivision.

3.23 (g) Prior authorization under this subdivision shall comply with section 62Q.184.

3.24 (h) Any step therapy protocol requirements established by the commissioner must comply  
3.25 with section 62Q.1841.