

SECOND REGULAR SESSION

SENATE BILL NO. 995

99TH GENERAL ASSEMBLY

INTRODUCED BY SENATOR SATER.

Read 1st time February 12, 2018, and ordered printed.

ADRIANE D. CROUSE, Secretary.

6373S.01I

AN ACT

To amend chapter 208, RSMo, by adding thereto one new section relating to an advisory council on rare diseases and personalized medicine.

Be it enacted by the General Assembly of the State of Missouri, as follows:

Section A. Chapter 208, RSMo, is amended by adding thereto one new
2 section, to be known as section 208.183, to read as follows:

**208.183. 1. There shall be established an "Advisory Council on
2 Rare Diseases and Personalized Medicine" within the MO HealthNet
3 division. The advisory council shall serve as an expert advisory
4 committee to the drug utilization review board, providing necessary
5 consultation to the board when the board makes recommendations or
6 determinations regarding beneficiary access to drugs or biological
7 products for rare diseases, or when the board itself determines that it
8 lacks the specific scientific, medical, or technical expertise necessary
9 for the proper performance of its responsibilities and such necessary
10 expertise can be provided by experts outside the board. "Beneficiary
11 access", as used in this section, shall mean developing prior
12 authorization and reauthorization criteria for a rare disease drug,
13 including placement on a preferred drug list or a formulary, as well as
14 payment, cost-sharing, drug utilization review, or medication therapy
15 management.**

**16 2. The advisory council on rare diseases and personalized
17 medicine shall be composed of the following health care professionals,
18 who shall be appointed by the director of the department of social
19 services:**

**20 (1) Two physicians affiliated with a public school of medicine
21 who are licensed and practicing in this state with experience**

22 researching, diagnosing, or treating rare diseases;

23 (2) Two physicians affiliated with private schools of medicine
24 headquartered in this state who are licensed and practicing in this
25 state with experience researching, diagnosing, or treating rare
26 diseases;

27 (3) A physician who holds a doctor of osteopathy degree, who is
28 active in medical practice, and who is affiliated with a school of
29 medicine in this state with experience researching, diagnosing, or
30 treating rare diseases;

31 (4) Two medical researchers from either academic research
32 institutions or medical research organizations in this state who have
33 received federal or foundation grant funding for rare disease research;

34 (5) A registered nurse or advanced practice registered nurse
35 licensed and practicing in this state with experience treating rare
36 diseases;

37 (6) A pharmacist practicing in a hospital in this state which has
38 a designated orphan disease center;

39 (7) A professor employed by a pharmacy program in this state
40 that is fully accredited by the Accreditation Council for Pharmacy
41 Education and who has advanced scientific or medical training in
42 orphan and rare disease treatments;

43 (8) One individual representing the rare disease community or
44 who is living with a rare disease;

45 (9) One member who represents a rare disease foundation;

46 (10) A representative from a rare disease center located within
47 one of the state's comprehensive pediatric hospitals;

48 (11) The chair of the joint committee on the life sciences or the
49 chair's designee; and

50 (12) The chairperson of the drug utilization review board, or the
51 chairperson's designee, who shall serve as an ex officio, nonvoting
52 member of the advisory council.

53 3. The director shall convene the first meeting of the advisory
54 council on rare diseases and personalized medicine no later than
55 February 28, 2019. Following the first meeting, the advisory council
56 shall meet upon the call of the chairperson of the drug utilization
57 review board or upon the request of a majority of the council members.

58 4. The drug utilization review board, when making

59 recommendations or determinations regarding beneficiary access to
60 drugs and biological products for rare diseases, as defined in the
61 federal Orphan Drug Act of 1983, P.L. 97-414, and drugs and biological
62 products that are approved by the U.S. Food and Drug Administration
63 and within the emerging fields of personalized medicine and
64 noninheritable gene editing therapeutics, shall request and consider
65 information from the advisory council on rare diseases and
66 personalized medicine.

67 5. The drug utilization review board shall seek the input of the
68 advisory council on rare diseases and personalized medicine to address
69 topics for consultation under this section including, but not limited to:

- 70 (1) Rare diseases;
- 71 (2) The severity of rare diseases;
- 72 (3) The unmet medical need associated with rare diseases;
- 73 (4) The impact of particular coverage, cost-sharing, tiering,
74 utilization management, prior authorization, medication therapy
75 management, or other Medicaid policies on access to rare disease
76 therapies;
- 77 (5) An assessment of the benefits and risks of therapies to treat
78 rare diseases;
- 79 (6) The impact of particular coverage, cost-sharing, tiering,
80 utilization management, prior authorization, medication therapy
81 management, or other policies on patients' adherence to the treatment
82 regimen prescribed or otherwise recommended by their physicians;
- 83 (7) Whether beneficiaries who need treatment from or a
84 consultation with a rare disease specialist have adequate access and,
85 if not, what factors are causing the limited access; and
- 86 (8) The demographics and the clinical description of patient
87 populations.

88 6. Nothing in this section shall be construed to create a legal
89 right for a consultation on any matter or to require the drug utilization
90 review board to meet with any particular expert or stakeholder.

91 7. Recommendations of the advisory council on rare diseases and
92 personalized medicine on an applicable treatment of a rare disease
93 shall be explained in writing to members of the drug utilization review
94 board during public hearings.

95 8. In cases of conflict where the drug utilization review board

96 makes a coverage decision that contradicts the recommendations of the
97 advisory council on rare diseases and personalized medicine, the board
98 shall clarify the reasoning behind such a decision in a publicly
99 available format including, but not limited to, published board minutes.

100 **9. For purposes of this section, a "rare disease drug" shall mean**
101 **a drug used to treat a rare medical condition, defined as any disease or**
102 **condition that affects fewer than two hundred thousand persons in the**
103 **United States, such as cystic fibrosis, hemophilia, and multiple**
104 **myeloma.**

105 **10. All members of the advisory council on rare diseases and**
106 **personalized medicine shall annually sign a conflict of interest**
107 **statement revealing economic or other relationships with entities that**
108 **could influence a member's decisions, and at least twenty percent of the**
109 **advisory council members shall not have a conflict of interest with**
110 **respect to any insurer, pharmaceutical benefits manager, or**
111 **pharmaceutical manufacturer.**

Bill ✓

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