

CERTIFICATION OF ENROLLMENT
ENGROSSED SECOND SUBSTITUTE HOUSE BILL 2575

59th Legislature
2006 Regular Session

Passed by the House March 6, 2006
Yeas 97 Nays 1

Speaker of the House of Representatives

Passed by the Senate March 3, 2006
Yeas 48 Nays 0

President of the Senate

Approved

Governor of the State of Washington

CERTIFICATE

I, Richard Nafziger, Chief Clerk of the House of Representatives of the State of Washington, do hereby certify that the attached is **ENGROSSED SECOND SUBSTITUTE HOUSE BILL 2575** as passed by the House of Representatives and the Senate on the dates hereon set forth.

Chief Clerk

FILED

**Secretary of State
State of Washington**

ENGROSSED SECOND SUBSTITUTE HOUSE BILL 2575

AS AMENDED BY THE SENATE

Passed Legislature - 2006 Regular Session

State of Washington 59th Legislature 2006 Regular Session

By House Committee on Appropriations (originally sponsored by Representatives Cody, Morrell and Moeller; by request of Governor Gregoire)

READ FIRST TIME 02/07/06.

1 AN ACT Relating to establishing a state health technology
2 assessment program; amending RCW 41.05.013; adding new sections to
3 chapter 70.14 RCW; and creating new sections.

4 BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF WASHINGTON:

5 NEW SECTION. **Sec. 1.** A new section is added to chapter 70.14 RCW
6 to read as follows:

7 DEFINITIONS. The definitions in this section apply throughout
8 sections 2 through 7 of this act unless the context clearly requires
9 otherwise.

10 (1) "Administrator" means the administrator of the Washington state
11 health care authority under chapter 41.05 RCW.

12 (2) "Advisory group" means a group established under section
13 4(2)(c) of this act.

14 (3) "Committee" means the health technology clinical committee
15 established under section 2 of this act.

16 (4) "Coverage determination" means a determination of the
17 circumstances, if any, under which a health technology will be included
18 as a covered benefit in a state purchased health care program.

1 (5) "Health technology" means medical and surgical devices and
2 procedures, medical equipment, and diagnostic tests. Health
3 technologies does not include prescription drugs governed by RCW
4 70.14.050.

5 (6) "Participating agency" means the department of social and
6 health services, the state health care authority, and the department of
7 labor and industries.

8 (7) "Reimbursement determination" means a determination to provide
9 or deny reimbursement for a health technology included as a covered
10 benefit in a specific circumstance for an individual patient who is
11 eligible to receive health care services from the state purchased
12 health care program making the determination.

13 NEW SECTION. **Sec. 2.** A new section is added to chapter 70.14 RCW
14 to read as follows:

15 HEALTH TECHNOLOGY COMMITTEE ESTABLISHED. (1) A health technology
16 clinical committee is established, to include the following eleven
17 members appointed by the administrator in consultation with
18 participating state agencies:

19 (a) Six practicing physicians licensed under chapter 18.57 or 18.71
20 RCW; and

21 (b) Five other practicing licensed health professionals who use
22 health technology in their scope of practice.

23 At least two members of the committee must have professional
24 experience treating women, children, elderly persons, and people with
25 diverse ethnic and racial backgrounds.

26 (2) Members of the committee:

27 (a) Shall not contract with or be employed by a health technology
28 manufacturer or a participating agency during their term or for
29 eighteen months before their appointment. As a condition of
30 appointment, each person shall agree to the terms and conditions
31 imposed by the administrator regarding conflicts of interest;

32 (b) Are immune from civil liability for any official acts performed
33 in good faith as members of the committee; and

34 (c) Shall be compensated for participation in the work of the
35 committee in accordance with a personal services contract to be
36 executed after appointment and before commencement of activities
37 related to the work of the committee.

1 (3) Meetings of the committee and any advisory group are subject to
2 chapter 42.30 RCW, the open public meetings act, including RCW
3 42.30.110(1)(1), which authorizes an executive session during a regular
4 or special meeting to consider proprietary or confidential nonpublished
5 information.

6 (4) Neither the committee nor any advisory group is an agency for
7 purposes of chapter 34.05 RCW.

8 (5) The health care authority shall provide administrative support
9 to the committee and any advisory group, and may adopt rules governing
10 their operation.

11 NEW SECTION. **Sec. 3.** A new section is added to chapter 70.14 RCW
12 to read as follows:

13 TECHNOLOGY SELECTION AND ASSESSMENT. (1) The administrator, in
14 consultation with participating agencies and the committee, shall
15 select the health technologies to be reviewed by the committee under
16 section 4 of this act. Up to six may be selected for review in the
17 first year after the effective date of this act, and up to eight may be
18 selected in the second year after the effective date of this act. In
19 making the selection, priority shall be given to any technology for
20 which:

21 (a) There are concerns about its safety, efficacy, or cost-
22 effectiveness, especially relative to existing alternatives, or
23 significant variations in its use;

24 (b) Actual or expected state expenditures are high, due to demand
25 for the technology, its cost, or both; and

26 (c) There is adequate evidence available to conduct the complete
27 review.

28 (2) A health technology for which the committee has made a
29 determination under section 4 of this act shall be considered for
30 rereview at least once every eighteen months, beginning the date the
31 determination is made. The administrator, in consultation with
32 participating agencies and the committee, shall select the technology
33 for rereview if he or she decides that evidence has since become
34 available that could change a previous determination. Upon rereview,
35 consideration shall be given only to evidence made available since the
36 previous determination.

1 (3) Pursuant to a petition submitted by an interested party, the
2 health technology clinical committee may select health technologies for
3 review that have not otherwise been selected by the administrator under
4 subsection (1) or (2) of this section.

5 (4) Upon the selection of a health technology for review, the
6 administrator shall contract for a systematic evidence-based assessment
7 of the technology's safety, efficacy, and cost-effectiveness. The
8 contract shall:

9 (a) Be with an evidence-based practice center designated as such by
10 the federal agency for health care research and quality, or other
11 appropriate entity;

12 (b) Require the assessment be initiated no sooner than thirty days
13 after notice of the selection of the health technology for review is
14 posted on the internet under section 7 of this act;

15 (c) Require, in addition to other information considered as part of
16 the assessment, consideration of: (i) Safety, health outcome, and cost
17 data submitted by a participating agency; and (ii) evidence submitted
18 by any interested party; and

19 (d) Require the assessment to: (i) Give the greatest weight to the
20 evidence determined, based on objective indicators, to be the most
21 valid and reliable, considering the nature and source of the evidence,
22 the empirical characteristic of the studies or trials upon which the
23 evidence is based, and the consistency of the outcome with comparable
24 studies; and (ii) take into account any unique impacts of the
25 technology on specific populations based upon factors such as sex, age,
26 ethnicity, race, or disability.

27 NEW SECTION. **Sec. 4.** A new section is added to chapter 70.14 RCW
28 to read as follows:

29 HEALTH TECHNOLOGY COMMITTEE DETERMINATIONS. (1) The committee
30 shall determine, for each health technology selected for review under
31 section 3 of this act: (a) The conditions, if any, under which the
32 health technology will be included as a covered benefit in health care
33 programs of participating agencies; and (b) if covered, the criteria
34 which the participating agency administering the program must use to
35 decide whether the technology is medically necessary, or proper and
36 necessary treatment.

1 (2) In making a determination under subsection (1) of this section,
2 the committee:

3 (a) Shall consider, in an open and transparent process, evidence
4 regarding the safety, efficacy, and cost-effectiveness of the
5 technology as set forth in the systematic assessment conducted under
6 section 3(4) of this act;

7 (b) Shall provide an opportunity for public comment; and

8 (c) May establish ad hoc temporary advisory groups if specialized
9 expertise is needed to review a particular health technology or group
10 of health technologies, or to seek input from enrollees or clients of
11 state purchased health care programs. Advisory group members are
12 immune from civil liability for any official act performed in good
13 faith as a member of the group. As a condition of appointment, each
14 person shall agree to the terms and conditions imposed by the
15 administrator regarding conflicts of interest.

16 (3) Determinations of the committee under subsection (1) of this
17 section shall be consistent with decisions made under the federal
18 medicare program and in expert treatment guidelines, including those
19 from specialty physician organizations and patient advocacy
20 organizations, unless the committee concludes, based on its review of
21 the systematic assessment, that substantial evidence regarding the
22 safety, efficacy, and cost-effectiveness of the technology supports a
23 contrary determination.

24 NEW SECTION. **Sec. 5.** A new section is added to chapter 70.14 RCW
25 to read as follows:

26 COMPLIANCE BY STATE AGENCIES. (1) A participating agency shall
27 comply with a determination of the committee under section 4 of this
28 act unless:

29 (a) The determination conflicts with an applicable federal statute
30 or regulation, or applicable state statute; or

31 (b) Reimbursement is provided under an agency policy regarding
32 experimental or investigational treatment, services under a clinical
33 investigation approved by an institutional review board, or health
34 technologies that have a humanitarian device exemption from the federal
35 food and drug administration.

36 (2) For a health technology not selected for review under section
37 3 of this act, a participating agency may use its existing statutory

1 and administrative authority to make coverage and reimbursement
2 determinations. Such determinations shall be shared among agencies,
3 with a goal of maximizing each agency's understanding of the basis for
4 the other's decisions and providing opportunities for agency
5 collaboration.

6 (3) A health technology not included as a covered benefit under a
7 state purchased health care program pursuant to a determination of the
8 health technology clinical committee under section 4 of this act, or
9 for which a condition of coverage established by the committee is not
10 met, shall not be subject to a determination in the case of an
11 individual patient as to whether it is medically necessary, or proper
12 and necessary treatment.

13 (4) Nothing in this act diminishes an individual's right under
14 existing law to appeal an action or decision of a participating agency
15 regarding a state purchased health care program. Appeals shall be
16 governed by state and federal law applicable to participating agency
17 decisions.

18 NEW SECTION. **Sec. 6.** A new section is added to chapter 70.14 RCW
19 to read as follows:

20 APPEAL PROCESS. The administrator shall establish an open,
21 independent, transparent, and timely process to enable patients,
22 providers, and other stakeholders to appeal the determinations of the
23 health technology clinical committee made under section 4 of this act.

24 NEW SECTION. **Sec. 7.** A new section is added to chapter 70.14 RCW
25 to read as follows:

26 PUBLIC NOTICE. (1) The administrator shall develop a centralized,
27 internet-based communication tool that provides, at a minimum:

28 (a) Notification when a health technology is selected for review
29 under section 3 of this act, indicating when the review will be
30 initiated and how an interested party may submit evidence, or provide
31 public comment, for consideration during the review;

32 (b) Notification of any determination made by the committee under
33 section 4(1) of this act, its effective date, and an explanation of the
34 basis for the determination; and

35 (c) Access to the systematic assessment completed under section

1 3(4) of this act, and reports completed under subsection (2) of this
2 section.

3 (2) Participating agencies shall develop methods to report on the
4 implementation of this section and sections 1 through 6 of this act
5 with respect to health care outcomes, frequency of exceptions, cost
6 outcomes, and other matters deemed appropriate by the administrator.

7 **Sec. 8.** RCW 41.05.013 and 2005 c 462 s 3 are each amended to read
8 as follows:

9 (1) The authority shall coordinate state agency efforts to develop
10 and implement uniform policies across state purchased health care
11 programs that will ensure prudent, cost-effective health services
12 purchasing, maximize efficiencies in administration of state purchased
13 health care programs, improve the quality of care provided through
14 state purchased health care programs, and reduce administrative burdens
15 on health care providers participating in state purchased health care
16 programs. The policies adopted should be based, to the extent
17 possible, upon the best available scientific and medical evidence and
18 shall endeavor to address:

19 (a) Methods of formal assessment, such as a health technology
20 assessment under sections 1 through 7 of this act. Consideration of
21 the best available scientific evidence does not preclude consideration
22 of experimental or investigational treatment or services under a
23 clinical investigation approved by an institutional review board;

24 (b) Monitoring of health outcomes, adverse events, quality, and
25 cost-effectiveness of health services;

26 (c) Development of a common definition of medical necessity; and

27 (d) Exploration of common strategies for disease management and
28 demand management programs, including asthma, diabetes, heart disease,
29 and similar common chronic diseases. Strategies to be explored include
30 individual asthma management plans. On January 1, 2007, and January 1,
31 2009, the authority shall issue a status report to the legislature
32 summarizing any results it attains in exploring and coordinating
33 strategies for asthma, diabetes, heart disease, and other chronic
34 diseases.

35 (2) The administrator may invite health care provider
36 organizations, carriers, other health care purchasers, and consumers to
37 participate in efforts undertaken under this section.

1 (3) For the purposes of this section "best available scientific and
2 medical evidence" means the best available clinical evidence derived
3 from systematic research.

4 NEW SECTION. **Sec. 9.** A new section is added to chapter 70.14 RCW
5 to read as follows:

6 Sections 1 through 7 of this act and RCW 41.05.013 do not apply to
7 state purchased health care services that are purchased from or through
8 health carriers as defined in RCW 48.43.005.

9 NEW SECTION. **Sec. 10.** Captions used in this act are not any part
10 of the law.

11 NEW SECTION. **Sec. 11.** If any part of this act is found to be in
12 conflict with federal requirements that are a prescribed condition to
13 the allocation of federal funds to the state, the conflicting part of
14 this act is inoperative solely to the extent of the conflict and with
15 respect to the agencies directly affected, and this finding does not
16 affect the operation of the remainder of this act in its application to
17 the agencies concerned. Rules adopted under this act must meet federal
18 requirements that are a necessary condition to the receipt of federal
19 funds by the state.

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