CERTIFICATION OF ENROLLMENT

ENGROSSED SUBSTITUTE HOUSE BILL 1414

Chapter 273, Laws of 2007

60th Legislature 2007 Regular Session

AMBULATORY SURGICAL FACILITIES--LICENSING

EFFECTIVE DATE: 07/01/09 - Except section 7, which becomes effective 07/22/07.

Passed by the House April 14, 2007 Yeas 91 Nays 2

FRANK CHOPP

Speaker of the House of Representatives

Passed by the Senate April 11, 2007 Yeas 48 Nays 1

BRAD OWEN

President of the Senate

Approved May 2, 2007, 11:01 a.m.

CERTIFICATE

I, Richard Nafziger, Chief Clerk of the House of Representatives of the State of Washington, do hereby certify that the attached is **ENGROSSED SUBSTITUTE HOUSE BILL** 1414 as passed by the House of Representatives and the Senate on the dates hereon set forth.

RICHARD NAFZIGER

Chief Clerk

FILED

May 3, 2007

Secretary of State State of Washington

CHRISTINE GREGOIRE

Governor of the State of Washington

ENGROSSED SUBSTITUTE HOUSE BILL 1414

AS AMENDED BY THE SENATE

Passed Legislature - 2007 Regular Session

State of Washington 60th Legislature 2007 Regular Session

By House Committee on Health Care & Wellness (originally sponsored by Representatives Cody, Green, Morrell, Moeller, Schual-Berke and Campbell)

READ FIRST TIME 02/12/07.

AN ACT Relating to licensing ambulatory surgical facilities; amending RCW 70.56.010, 18.130.070, 18.71.0195, 18.71.017, 18.57.005, and 18.22.015; reenacting and amending RCW 43.70.510, 70.41.200, and 42.56.360; adding a new chapter to Title 70 RCW; creating new sections; prescribing penalties; and providing an effective date.

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

7 **Sec. 1.** The definitions in this section apply NEW SECTION. 8 throughout this chapter unless the context clearly requires otherwise. 9 (1) "Ambulatory surgical facility" means any distinct entity that 10 operates for the primary purpose of providing specialty or multispecialty outpatient surgical services in which patients are 11 12 admitted to and discharged from the facility within twenty-four hours and do not require inpatient hospitalization, whether or not the 13 14 facility is certified under Title XVIII of the federal social security 15 act.

16 (2) "Department" means the department of health.

17 (3) "General anesthesia" means a state of unconsciousness18 intentionally produced by anesthetic agents, with absence of pain

1 sensation over the entire body, in which the patient is without 2 protective reflexes and is unable to maintain an airway.

3 (4) "Person" means an individual, firm, partnership, corporation,
4 company, association, joint stock association, and the legal successor
5 thereof.

6 (5) "Practitioner" means any physician or surgeon licensed under 7 chapter 18.71 RCW, an osteopathic physician or surgeon licensed under 8 chapter 18.57 RCW, or a podiatric physician or surgeon licensed under 9 chapter 18.22 RCW.

10

(6) "Secretary" means the secretary of health.

11 (7) "Surgical services" means invasive medical procedures that:

(a) Utilize a knife, laser, cautery, cryogenics, or chemicals; and
(b) Remove, correct, or facilitate the diagnosis or cure of a
disease, process, or injury through that branch of medicine that treats
diseases, injuries, and deformities by manual or operative methods by
a practitioner.

17 <u>NEW SECTION.</u> Sec. 2. The secretary shall:

18 (1) Issue a license to any ambulatory surgical facility that:

19 (a) Submits payment of the fee established in section 7 of this 20 act;

21 (b) Submits a completed application that demonstrates the ability 22 to comply with the standards established for operating and maintaining 23 an ambulatory surgical facility in statute and rule. An ambulatory 24 surgical facility shall be deemed to have met the standards if it submits proof of certification as a medicare ambulatory surgical 25 26 facility or accreditation by an organization that the secretary has 27 determined to have substantially equivalent standards to those of the 28 department; and

29 (c) Successfully completes the survey requirements established in 30 section 11 of this act;

31 (2) Develop an application form for applicants for a license to 32 operate an ambulatory surgical facility;

(3) Initiate investigations and enforcement actions for complaints
 or other information regarding failure to comply with this chapter or
 the standards and rules adopted under this chapter;

36 (4) Conduct surveys of facilities, including reviews of medical

1 records and documents required to be maintained under this chapter or 2 rules adopted under this chapter;

3 (5) By March 1, 2008, determine which accreditation organizations 4 have substantially equivalent standards for purposes of deeming 5 specific licensing requirements required in statute and rule as having 6 met the state's standards; and

7

(6) Adopt any rules necessary to implement this chapter.

8 <u>NEW SECTION.</u> Sec. 3. Except as provided in section 4 of this act, after June 30, 2009, no person or governmental unit of the state of 9 10 Washington, acting separately or jointly with any other person or 11 governmental unit, shall establish, maintain, or conduct an ambulatory 12 surgical facility in this state or advertise by using the term "ambulatory surgical facility," "day surgery center," "licensed 13 surgical center," or other words conveying similar meaning without a 14 15 license issued by the department under this chapter.

16 <u>NEW SECTION.</u> Sec. 4. Nothing in this chapter:

(1) Applies to an ambulatory surgical facility that is maintainedand operated by a hospital licensed under chapter 70.41 RCW;

19 (2) Applies to an office maintained for the practice of dentistry;

20 (3) Applies to outpatient specialty or multispecialty surgical 21 services routinely and customarily performed in the office of a 22 practitioner in an individual or group practice that do not require 23 general anesthesia; or

24 (4) Limits an ambulatory surgical facility to performing only25 surgical services.

26 <u>NEW SECTION.</u> Sec. 5. (1) An applicant for a license to operate an 27 ambulatory surgical facility must demonstrate the ability to comply 28 with the standards established for operating and maintaining an 29 ambulatory surgical facility in statute and rule, including:

30 (a) Submitting a written application to the department providing
 31 all necessary information on a form provided by the department,
 32 including a list of surgical specialties offered;

33 (b) Submitting building plans for review and approval by the 34 department for new construction, alterations other than minor alterations, and additions to existing facilities, prior to obtaining
 a license and occupying the building;

3 (c) Demonstrating the ability to comply with this chapter and any
4 rules adopted under this chapter;

(d) Cooperating with the department during on-site surveys prior to
obtaining an initial license or renewing an existing license;

7 (e) Providing such proof as the department may require concerning 8 the ownership and management of the ambulatory surgical facility, 9 including information about the organization and governance of the 10 facility and the identity of the applicant, officers, directors, 11 partners, managing employees, or owners of ten percent or more of the 12 applicant's assets;

13 (f) Submitting proof of operation of a coordinated quality 14 improvement program in accordance with section 9 of this act;

(g) Submitting a copy of the facility safety and emergency training program established under section 6 of this act;

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(h) Paying any fees established under section 7 of this act; and

18 (i) Providing any other information that the department may 19 reasonably require.

(2) A license is valid for three years, after which an ambulatory 20 surgical facility must submit an application for renewal of license 21 22 upon forms provided by the department and the renewal fee as established in section 7 of this act. The applicant must demonstrate 23 24 the ability to comply with the standards established for operating and 25 maintaining an ambulatory surgical facility in statutes, standards, and rules. The applicant must submit the license renewal document no later 26 27 than thirty days prior to the date of expiration of the license.

(3) The applicant may demonstrate compliance with any of the requirements of subsection (1) of this section by providing satisfactory documentation to the secretary that it has met the standards of an accreditation organization or federal agency that the secretary has determined to have substantially equivalent standards as the statutes and rules of this state.

34 <u>NEW SECTION.</u> Sec. 6. An ambulatory surgical facility shall have 35 a facility safety and emergency training program. The program shall 36 include:

1 (1) On-site equipment, medication, and trained personnel to 2 facilitate handling of services sought or provided and to facilitate 3 the management of any medical emergency that may arise in connection 4 with services sought or provided;

5 (2) Written transfer agreements with local hospitals licensed under 6 chapter 70.41 RCW, approved by the ambulatory surgical facility's 7 medical staff; and

8 (3) A procedural plan for handling medical emergencies that shall
9 be available for review during surveys and inspections.

10 <u>NEW SECTION.</u> Sec. 7. The department of health shall convene a 11 group of interested stakeholders to identify relevant regulatory issues 12 related to the implementation of this act, including a reasonable fee 13 schedule for licenses and renewal licenses. The group shall report to 14 the department on their recommendations no later than December 15, 15 2007.

16 <u>NEW SECTION.</u> Sec. 8. (1) The secretary may deny, suspend, or 17 revoke the license of any ambulatory surgical facility in any case in 18 which he or she finds the applicant or registered entity knowingly made 19 a false statement of material fact in the application for the license 20 or any supporting data in any record required by this chapter or matter 21 under investigation by the department.

22 (2) The secretary shall investigate complaints concerning operation 23 of an ambulatory surgical facility without a license. The secretary may issue a notice of intention to issue a cease and desist order to 24 25 any person whom the secretary has reason to believe is engaged in the unlicensed operation of an ambulatory surgical facility. 26 If the secretary makes a written finding of fact that the public interest will 27 be irreparably harmed by delay in issuing an order, the secretary may 28 issue a temporary cease and desist order. The person receiving a 29 30 temporary cease and desist order shall be provided an opportunity for a prompt hearing. The temporary cease and desist order shall remain in 31 effect until further order of the secretary. Any person operating an 32 ambulatory surgical facility under this chapter without a license is 33 34 quilty of a misdemeanor, and each day of operation of an unlicensed 35 ambulatory surgical facility constitutes a separate offense.

(3) The secretary is authorized to deny, suspend, revoke, or modify a license or provisional license in any case in which it finds that there has been a failure or refusal to comply with the requirements of this chapter or the standards or rules adopted under this chapter. RCW 43.70.115 governs notice of a license denial, revocation, suspension, or modification and provides the right to an adjudicative proceeding.

7 (4) Pursuant to chapter 34.05 RCW, the secretary may assess
8 monetary penalties of a civil nature not to exceed one thousand dollars
9 per violation.

10 <u>NEW SECTION.</u> Sec. 9. (1) Every ambulatory surgical facility shall 11 maintain a coordinated quality improvement program for the improvement 12 of the quality of health care services rendered to patients and the 13 identification and prevention of medical malpractice. The program 14 shall include at least the following:

15 (a) The establishment of a quality improvement committee with the 16 responsibility to review the services rendered in the ambulatory 17 surgical facility, both retrospectively and prospectively, in order to improve the quality of medical care of patients and to prevent medical 18 malpractice. The committee shall oversee and coordinate the quality 19 improvement and medical malpractice prevention program and shall ensure 20 21 that information gathered pursuant to the program is used to review and 22 to revise the policies and procedures of the ambulatory surgical facility; 23

(b) A medical staff privileges sanction procedure through which credentials, physical and mental capacity, and competence in delivering health care services are periodically reviewed as part of an evaluation of staff privileges;

(c) The periodic review of the credentials, physical and mental capacity, and competence in delivering health care services of all persons who are employed or associated with the ambulatory surgical facility;

(d) A procedure for the prompt resolution of grievances by patients
 or their representatives related to accidents, injuries, treatment, and
 other events that may result in claims of medical malpractice;

(e) The maintenance and continuous collection of information
 concerning the ambulatory surgical facility's experience with negative
 health care outcomes and incidents injurious to patients, patient

grievances, professional liability premiums, settlements, awards, costs incurred by the ambulatory surgical facility for patient injury prevention, and safety improvement activities;

4 (f) The maintenance of relevant and appropriate information
5 gathered pursuant to (a) through (e) of this subsection concerning
6 individual practitioners within the practitioner's personnel or
7 credential file maintained by the ambulatory surgical facility;

8 (g) Education programs dealing with quality improvement, patient 9 safety, medication errors, injury prevention, staff responsibility to 10 report professional misconduct, the legal aspects of patient care, 11 improved communication with patients, and causes of malpractice claims 12 for staff personnel engaged in patient care activities; and

(h) Policies to ensure compliance with the reporting requirementsof this section.

(2) Any person who, in substantial good faith, provides information 15 to further the purposes of the quality improvement and medical 16 17 malpractice prevention program or who, in substantial good faith, participates on the quality improvement committee is not subject to an 18 action for civil damages or other relief as a result of such activity. 19 Any person or entity participating in a coordinated quality improvement 20 21 program that, in substantial good faith, shares information or 22 documents with one or more other programs, committees, or boards under subsection (8) of this section is not subject to an action for civil 23 24 damages or other relief as a result of the activity. For the purposes 25 of this section, sharing information is presumed to be in substantial good faith. However, the presumption may be rebutted upon a showing of 26 27 clear, cogent, and convincing evidence that the information shared was knowingly false or deliberately misleading. 28

(3) Information and documents, including complaints and incident 29 reports, created specifically for, and collected and maintained by, a 30 31 quality improvement committee are not subject to review or disclosure, 32 except as provided in this section, or discovery or introduction into evidence in any civil action, and no person who was in attendance at a 33 meeting of such committee or who participated in the creation, 34 collection, or maintenance of information or documents specifically for 35 the committee shall be permitted or required to testify in any civil 36 37 action as to the content of such proceedings or the documents and 38 information prepared specifically for the committee. This subsection

does not preclude: (a) In any civil action, the discovery of the 1 2 identity of persons involved in the medical care that is the basis of the civil action whose involvement was independent of any quality 3 improvement activity; (b) in any civil action, the testimony of any 4 person concerning the facts which form the basis for the institution of 5 such proceedings of which the person had personal knowledge acquired 6 7 independently of such proceedings; (c) in any civil action by a health care provider regarding the restriction or revocation of that 8 individual's clinical or staff privileges, introduction into evidence 9 10 of information collected and maintained by quality improvement committees regarding such health care provider; (d) in any civil 11 12 action, disclosure of the fact that staff privileges were terminated or 13 restricted, including the specific restrictions imposed, if any, and 14 the reasons for the restrictions; or (e) in any civil action, discovery and introduction into evidence of the patient's medical records 15 required by rule of the department to be made regarding the care and 16 17 treatment received.

18 (4) Each quality improvement committee shall, on at least a 19 semiannual basis, report to the management of the ambulatory surgical 20 facility, as identified in the facility's application, in which the 21 committee is located. The report shall review the quality improvement 22 activities conducted by the committee, and any actions taken as a 23 result of those activities.

(5) The department shall adopt such rules as are deemed appropriateto effectuate the purposes of this section.

The medical quality assurance commission, the board of 26 (6) 27 osteopathic medicine and surgery, or the podiatric medical board, as appropriate, may review and audit the records of committee decisions in 28 which a practitioner's privileges are terminated or restricted. Each 29 ambulatory surgical facility shall produce and make accessible to the 30 commission or board the appropriate records and otherwise facilitate 31 32 the review and audit. Information so gained is not subject to the discovery process and confidentiality shall be respected as required by 33 subsection (3) of this section. Failure of an ambulatory surgical 34 35 facility to comply with this subsection is punishable by a civil penalty not to exceed two hundred fifty dollars. 36

37 (7) The department and any accrediting organization may review and38 audit the records of a quality improvement committee or peer review

1 committee in connection with their inspection and review of the 2 ambulatory surgical facility. Information so obtained is not subject 3 to the discovery process, and confidentiality shall be respected as 4 required by subsection (3) of this section. Each ambulatory surgical 5 facility shall produce and make accessible to the department the 6 appropriate records and otherwise facilitate the review and audit.

7 (8) A coordinated quality improvement program may share information and documents, including complaints and incident reports, created 8 specifically for, and collected and maintained by, a 9 quality improvement committee or a peer review committee under RCW 4.24.250 10 with one or more other coordinated quality improvement programs 11 12 maintained in accordance with this section or RCW 43.70.510 or 13 70.41.200, a quality assurance committee maintained in accordance with RCW 18.20.390 or 74.42.640, or a peer review committee under RCW 14 4.24.250, for the improvement of the quality of health care services 15 rendered to patients and the identification and prevention of medical 16 17 malpractice. The privacy protections of chapter 70.02 RCW and the federal health insurance portability and accountability act of 1996 and 18 its implementing regulations apply to the sharing of individually 19 identifiable patient information held by a coordinated quality 20 21 improvement program. Any rules necessary to implement this section 22 shall meet the requirements of applicable federal and state privacy Information and documents disclosed by one coordinated quality 23 laws. 24 improvement program to another coordinated quality improvement program 25 or a peer review committee under RCW 4.24.250 and any information and 26 documents created or maintained as a result of the sharing of 27 information and documents are not subject to the discovery process and confidentiality shall be respected as required by subsection (3) of 28 this section, RCW 18.20.390 (6) and (8), 70.41.200(3), 74.42.640 (7) 29 and (9), and 4.24.250. 30

31 (9) An ambulatory surgical facility that participates in a 32 coordinated quality improvement program under RCW 43.70.510 shall be 33 deemed to have met the requirements of this section.

34 (10) Violation of this section shall not be considered negligence 35 per se.

36 <u>NEW SECTION.</u> Sec. 10. The department shall establish and adopt 37 such minimum standards and rules pertaining to the construction,

maintenance, and operation of ambulatory surgical facilities and 1 2 rescind, amend, or modify such rules, as are necessary in the public interest, and particularly for the establishment and maintenance of 3 standards of patient care required for the safe and adequate care and 4 5 treatment of patients. In establishing the format and content of these standards and rules, the department shall give consideration to 6 7 maintaining consistency with such minimum standards and rules applicable to ambulatory surgical facilities in the survey standards of 8 accrediting organizations or federal agencies that the secretary has 9 10 determined to have substantially equivalent standards as the statutes and rules of this state. 11

12 <u>NEW SECTION.</u> Sec. 11. (1) The department shall make or cause to be made a survey of all ambulatory surgical facilities every eighteen 13 months. Every survey of an ambulatory surgical facility may include an 14 15 inspection of every part of the surgical facility. The department may 16 make an examination of all phases of the ambulatory surgical facility operation necessary to determine compliance with all applicable 17 statutes, rules, and regulations. In the event that the department is 18 unable to make a survey or cause a survey to be made during the three 19 20 years of the term of the license, the license of the ambulatory 21 surgical facility shall remain in effect until the state conducts a survey or a substitute survey is performed if the ambulatory surgical 22 23 facility is in compliance with all other licensing requirements.

24 (2) An ambulatory surgical facility shall be deemed to have met the survey standards of this section if it submits proof of certification 25 26 as a medicare ambulatory surgical facility or accreditation by an 27 organization that the secretary has determined to have substantially equivalent survey standards to those of the department. 28 A survey 29 performed pursuant to medicare certification or by an approved accrediting organization may substitute for a survey by the department 30 if: 31

32 (a) The ambulatory surgical facility has satisfactorily completed33 a survey by the department in the previous eighteen months; and

34 (b) Within thirty days of learning the result of a survey, the 35 ambulatory surgical facility provides the department with documentary 36 evidence that the ambulatory surgical facility has been certified or 37 accredited as a result of a survey and the date of the survey. (3) Ambulatory surgical facilities shall make the written reports
 of surveys conducted pursuant to medicare certification procedures or
 by an approved accrediting organization available to department
 surveyors during any department surveys, upon request.

5 <u>NEW SECTION.</u> Sec. 12. The department shall require ambulatory 6 surgical facilities to submit data related to the quality of patient 7 care for review by the department. The data shall be submitted every eighteen months. The department shall consider the reporting standards 8 9 of other public and private organizations that measure quality in order to maintain consistency in reporting and minimize the burden on the 10 11 ambulatory surgical facility. The department shall review the data to 12 determine the maintenance of quality patient care at the facility. If the department determines that the care offered at the facility may 13 present a risk to the health and safety of patients, the department may 14 15 conduct an inspection of the facility and initiate appropriate actions 16 to protect the public. Information submitted to the department 17 pursuant to this section shall be exempt from disclosure under chapter 42.56 RCW. 18

19 NEW SECTION. Sec. 13. (1) The chief administrator or executive officer of an ambulatory surgical facility shall report to the 20 department when the practice of a health care provider licensed by a 21 22 disciplining authority under RCW 18.130.040 is restricted, suspended, 23 limited, or terminated based upon a conviction, determination, or 24 finding by the ambulatory surgical facility that the provider has 25 committed an action defined as unprofessional conduct under RCW 18.130.180. The chief administrator or executive officer shall also 26 report any voluntary restriction or termination of the practice of a 27 health care provider licensed by a disciplining authority under RCW 28 18.130.040 while the provider is under investigation or the subject of 29 30 a proceeding by the ambulatory surgical facility regarding unprofessional conduct, or in return for the ambulatory surgical 31 32 facility not conducting such an investigation or proceeding or not taking action. The department shall forward the report to the 33 34 appropriate disciplining authority.

35 (2) Reports made under subsection (1) of this section must be made
 36 within fifteen days of the date of: (a) A conviction, determination,

or finding by the ambulatory surgical facility that the health care provider has committed an action defined as unprofessional conduct under RCW 18.130.180; or (b) acceptance by the ambulatory surgical facility of the voluntary restriction or termination of the practice of a health care provider, including his or her voluntary resignation, while under investigation or the subject of proceedings regarding unprofessional conduct under RCW 18.130.180.

8 (3) Failure of an ambulatory surgical facility to comply with this 9 section is punishable by a civil penalty not to exceed two hundred 10 fifty dollars.

(4) An ambulatory surgical facility, its chief administrator, or 11 its executive officer who files a report under this section is immune 12 13 from suit, whether direct or derivative, in any civil action related to 14 the filing or contents of the report, unless the conviction, determination, or finding on which the report and its content are based 15 is proven to not have been made in good faith. The prevailing party in 16 17 any action brought alleging that the conviction, determination, finding, or report was not made in good faith is entitled to recover 18 the costs of litigation, including reasonable attorneys' fees. 19

20 (5) The department shall forward reports made under subsection (1) 21 of this section to the appropriate disciplining authority designated 22 under Title 18 RCW within fifteen days of the date the report is 23 received by the department. The department shall notify an ambulatory 24 surgical facility that has made a report under subsection (1) of this 25 section of the results of the disciplining authority's case disposition decision within fifteen days after the case disposition. 26 Case 27 disposition is the decision whether to issue a statement of charges, take informal action, or close the complaint without action against a 28 In its biennial report to the legislature under RCW 29 provider. 18.130.310, the department shall specifically identify the case 30 dispositions of reports made by ambulatory surgical facilities under 31 32 subsection (1) of this section.

33 <u>NEW SECTION.</u> Sec. 14. Each ambulatory surgical facility shall 34 keep written records of decisions to restrict or terminate privileges 35 of practitioners. Copies of such records shall be made available to 36 the medical quality assurance commission, the board of osteopathic 37 medicine and surgery, or the podiatric medical board, within thirty

1 days of a request, and all information so gained remains confidential 2 in accordance with sections 9 and 13 of this act and is protected from 3 the discovery process. Failure of an ambulatory surgical facility to 4 comply with this section is punishable by a civil penalty not to exceed 5 two hundred fifty dollars.

6 <u>NEW SECTION.</u> **Sec. 15.** (1) Prior to granting or renewing clinical 7 privileges or association of any practitioner or hiring a practitioner, 8 an ambulatory surgical facility approved pursuant to this chapter shall 9 request from the practitioner and the practitioner shall provide the 10 following information:

(a) The name of any hospital, ambulatory surgical facility, or other facility with or at which the practitioner had or has any association, employment, privileges, or practice;

(b) If such association, employment, privilege, or practice wasdiscontinued, the reasons for its discontinuation;

16 (c) Any pending professional medical misconduct proceedings or any 17 pending medical malpractice actions in this state or another state, the 18 substance of the allegations in the proceedings or actions, and any 19 additional information concerning the proceedings or actions as the 20 practitioner deems appropriate;

(d) The substance of the findings in the actions or proceedings and any additional information concerning the actions or proceedings as the practitioner deems appropriate;

(e) A waiver by the practitioner of any confidentiality provisions
 concerning the information required to be provided to ambulatory
 surgical facilities pursuant to this subsection; and

(f) A verification by the practitioner that the informationprovided by the practitioner is accurate and complete.

(2) Prior to granting privileges or association to any practitioner or hiring a practitioner, an ambulatory surgical facility approved under this chapter shall request from any hospital or ambulatory surgical facility with or at which the practitioner had or has privileges, was associated, or was employed, the following information concerning the practitioner:

(a) Any pending professional medical misconduct proceedings or any
 pending medical malpractice actions, in this state or another state;

(b) Any judgment or settlement of a medical malpractice action and
 any finding of professional misconduct in this state or another state
 by a licensing or disciplinary board; and

4 (c) Any information required to be reported by hospitals or 5 ambulatory surgical facilities pursuant to RCW 18.130.070.

6 (3) The medical quality assurance commission, board of osteopathic 7 medicine and surgery, podiatric medical board, or dental quality 8 assurance commission, as appropriate, shall be advised within thirty 9 days of the name of any practitioner denied staff privileges, 10 association, or employment on the basis of adverse findings under 11 subsection (1) of this section.

(4) A hospital, ambulatory surgical facility, or other facility 12 13 that receives a request for information from another hospital, 14 ambulatory surgical facility, or other facility pursuant to subsections (1) and (2) of this section shall provide such information concerning 15 16 the physician in question to the extent such information is known to 17 the hospital, ambulatory surgical facility, or other facility receiving such a request, including the reasons for suspension, termination, or 18 curtailment of employment or privileges at the hospital, ambulatory 19 20 surgical facility, or facility. A hospital, ambulatory surgical 21 facility, other facility, or other person providing such information in 22 good faith is not liable in any civil action for the release of such 23 information.

24 (5) Information and documents, including complaints and incident 25 reports, created specifically for, and collected and maintained by, a quality improvement committee are not subject to discovery or 26 27 introduction into evidence in any civil action, and no person who was in attendance at a meeting of such committee or who participated in the 28 creation, collection, or maintenance of information or documents 29 specifically for the committee shall be permitted or required to 30 testify in any civil action as to the content of such proceedings or 31 32 the documents and information prepared specifically for the committee. This subsection does not preclude: (a) In any civil action, the 33 discovery of the identity of persons involved in the medical care that 34 is the basis of the civil action whose involvement was independent of 35 any quality improvement activity; (b) in any civil action, the 36 37 testimony of any person concerning the facts which form the basis for 38 the institution of such proceedings of which the person had personal

knowledge acquired independently of such proceedings; (c) in any civil 1 2 action by a health care provider regarding the restriction or individual's clinical or staff revocation of that 3 privileges, introduction into evidence information collected and maintained by 4 5 quality improvement committees regarding such health care provider; (d) in any civil action, disclosure of the fact that staff privileges were б 7 terminated or restricted, including the specific restrictions imposed, if any, and the reasons for the restrictions; or (e) in any civil 8 action, discovery and introduction into evidence of the patient's 9 10 medical records required by rule of the department to be made regarding the care and treatment received. 11

12 (6) Ambulatory surgical facilities shall be granted access to 13 information held by the medical quality assurance commission, board of 14 osteopathic medicine and surgery, or podiatric medical board pertinent 15 to decisions of the ambulatory surgical facility regarding 16 credentialing and recredentialing of practitioners.

17 (7) Violation of this section shall not be considered negligence18 per se.

NEW SECTION. Sec. 16. Ambulatory surgical facilities shall have 19 in place policies to assure that, when appropriate, information about 20 21 unanticipated outcomes is provided to patients or their families or any surrogate decision makers identified pursuant to RCW 7.70.065. 22 23 Notifications of unanticipated outcomes under this section do not 24 constitute an acknowledgement or admission of liability, nor may the fact of notification, the content disclosed, or any and all statements, 25 26 affirmations, gestures, or conduct expressing apology be introduced as evidence in a civil action. 27

28 <u>NEW SECTION.</u> Sec. 17. Every ambulatory surgical facility shall 29 post in conspicuous locations a notice of the department's ambulatory 30 surgical facility complaint toll-free telephone number. The form of 31 the notice shall be approved by the department.

32 <u>NEW SECTION.</u> Sec. 18. Information received by the department 33 through filed reports, inspection, or as otherwise authorized under 34 this chapter may be disclosed publicly, as permitted under chapter 35 42.56 RCW, subject to the following provisions:

1 (1) Licensing inspections, or complaint investigations regardless 2 of findings, shall, as requested, be disclosed no sooner than three 3 business days after the ambulatory surgical facility has received the 4 resulting assessment report;

5 (2) Information regarding administrative action against the license 6 shall, as requested, be disclosed after the ambulatory surgical 7 facility has received the documents initiating the administrative 8 action;

9 (3) Information about complaints that did not warrant an 10 investigation shall not be disclosed except to notify the ambulatory 11 surgical facility and the complainant that the complaint did not 12 warrant an investigation; and

13 (4) Information disclosed under this section shall not disclose 14 individual names.

<u>NEW SECTION.</u> Sec. 19. The ambulatory surgical facility account is 15 16 created in the custody of the state treasurer. All receipts from fees 17 and penalties imposed under this chapter must be deposited into the Expenditures from the account may be used only for 18 account. administration of this chapter. Only the secretary or the secretary's 19 20 designee may authorize expenditures from the account. The account is 21 subject to allotment procedures under chapter 43.88 RCW, but an appropriation is not required for expenditures. 22

23 **Sec. 20.** RCW 70.56.010 and 2006 c 8 s 105 are each amended to read 24 as follows:

The definitions in this section apply throughout this chapter unless the context clearly requires otherwise.

(1) "Adverse health event" or "adverse event" means the list of serious reportable events adopted by the national quality forum in 2002, in its consensus report on serious reportable events in health care. The department shall update the list, through adoption of rules, as subsequent changes are made by the national quality forum. The term does not include an incident.

(2) "Ambulatory surgical facility" means ((any distinct entity that
 operates exclusively for the purpose of providing surgical services to
 patients not requiring hospitalization, whether or not the facility is

certified under Title XVIII of the federal social security act)) a 1 facility licensed under chapter 70. -- RCW (sections 1 through 19 of 2 this act). 3

(3) "Childbirth center" means a facility licensed under chapter 4 5 18.46 RCW.

(4) "Correctional medical facility" means a part or unit of a 6 7 correctional facility operated by the department of corrections under chapter 72.10 RCW that provides medical services for lengths of stay in 8 9 excess of twenty-four hours to offenders.

10

(5) "Department" means the department of health.

(6) "Health care worker" means an employee, independent contractor, 11 licensee, or other individual who is directly involved in the delivery 12 13 of health services in a medical facility.

14

(7) "Hospital" means a facility licensed under chapter 70.41 RCW.

(8) "Incident" means an event, occurrence, or situation involving 15 16 the clinical care of a patient in a medical facility that:

17 (a) Results in unanticipated injury to a patient that is not related to the natural course of the patient's illness or underlying 18 condition and does not constitute an adverse event; or 19

(b) Could have injured the patient but did not either cause an 20 21 unanticipated injury or require the delivery of additional health care 22 services to the patient.

23

"Incident" does not include an adverse event.

24 (9) "Independent entity" means that entity that the department of 25 health contracts with under RCW 70.56.040 to receive notifications and reports of adverse events and incidents, and carry out the activities 26 27 specified in RCW 70.56.040.

(10) "Medical facility" means a childbirth center, hospital, 28 psychiatric hospital, or correctional medical facility. An ambulatory 29 surgical facility shall be considered a medical facility for purposes 30 31 of this chapter upon the effective date of any requirement for state 32 registration or licensure of ambulatory surgical facilities.

(11) "Psychiatric hospital" means a hospital facility licensed as 33 a psychiatric hospital under chapter 71.12 RCW. 34

sec. 21. RCW 43.70.510 and 2006 c 8 s 113, 2005 c 291 s 2, 2005 c 35 36 274 s 302, and 2005 c 33 s 6 are each reenacted and amended to read as 37 follows:

(1)(a) Health care institutions and medical facilities, other than 1 2 hospitals, that are licensed by the department, professional societies or organizations, health care service contractors, health maintenance 3 organizations, health carriers approved pursuant to chapter 48.43 RCW, 4 and any other person or entity providing health care coverage under 5 chapter 48.42 RCW that is subject to the jurisdiction and regulation of б 7 any state agency or any subdivision thereof may maintain a coordinated quality improvement program for the improvement of the quality of 8 health care services rendered to patients and the identification and 9 10 prevention of medical malpractice as set forth in RCW 70.41.200.

(b) All such programs shall comply with the requirements of RCW 11 12 70.41.200(1) (a), (c), (d), (e), (f), (g), and (h) as modified to 13 reflect the structural organization of the institution, facility, 14 professional societies organizations, health or care service contractors, health maintenance organizations, health carriers, or any 15 other person or entity providing health care coverage under chapter 16 17 48.42 RCW that is subject to the jurisdiction and regulation of any state agency or any subdivision thereof, unless an alternative quality 18 improvement program substantially equivalent to RCW 70.41.200(1)(a) is 19 developed. All such programs, whether complying with the requirement 20 21 set forth in RCW 70.41.200(1)(a) or in the form of an alternative 22 program, must be approved by the department before the discovery limitations provided in subsections (3) and (4) of this section and the 23 24 exemption under RCW 42.56.360(1)(c) and subsection (5) of this section 25 shall apply. In reviewing plans submitted by licensed entities that are associated with physicians' offices, the department shall ensure 26 27 that the exemption under RCW 42.56.360(1)(c) and the discovery limitations of this section are applied only to information and 28 documents related specifically to quality improvement activities 29 undertaken by the licensed entity. 30

(2) Health care provider groups of five or more providers may 31 32 maintain a coordinated quality improvement program for the improvement of the quality of health care services rendered to patients and the 33 identification and prevention of medical malpractice as set forth in 34 RCW 70.41.200. For purposes of this section, a health care provider 35 36 group may be a consortium of providers consisting of five or more 37 providers in total. All such programs shall comply with the 38 requirements of RCW 70.41.200(1) (a), (c), (d), (e), (f), (g), and (h)

as modified to reflect the structural organization of the health care provider group. All such programs must be approved by the department before the discovery limitations provided in subsections (3) and (4) of this section and the exemption under RCW 42.56.360(1)(c) and subsection (5) of this section shall apply.

(3) Any person who, in substantial good faith, provides information 6 7 to further the purposes of the quality improvement and medical malpractice prevention program or who, in substantial good faith, 8 participates on the quality improvement committee shall not be subject 9 10 to an action for civil damages or other relief as a result of such activity. Any person or entity participating in a coordinated quality 11 12 improvement program that, in substantial good faith, shares information or documents with one or more other programs, committees, or boards 13 under subsection (6) of this section is not subject to an action for 14 civil damages or other relief as a result of the activity or its 15 consequences. For the purposes of this section, sharing information is 16 17 presumed to be in substantial good faith. However, the presumption may be rebutted upon a showing of clear, cogent, and convincing evidence 18 that the information shared was knowingly false or deliberately 19 misleading. 20

21 (4) Information and documents, including complaints and incident reports, created specifically for, and collected and maintained by, a 22 quality improvement committee are not subject to review or disclosure, 23 24 except as provided in this section, or discovery or introduction into 25 evidence in any civil action, and no person who was in attendance at a meeting of such committee or who participated in the creation, 26 27 collection, or maintenance of information or documents specifically for the committee shall be permitted or required to testify in any civil 28 action as to the content of such proceedings or the documents and 29 information prepared specifically for the committee. This subsection 30 (a) In any civil action, the discovery of the 31 does not preclude: 32 identity of persons involved in the medical care that is the basis of the civil action whose involvement was independent of any quality 33 improvement activity; (b) in any civil action, the testimony of any 34 person concerning the facts that form the basis for the institution of 35 such proceedings of which the person had personal knowledge acquired 36 37 independently of such proceedings; (c) in any civil action by a health care provider regarding the restriction or revocation of that 38

individual's clinical or staff privileges, introduction into evidence 1 2 information collected and maintained by quality improvement committees regarding such health care provider; (d) in any civil action 3 challenging the termination of a contract by a state agency with any 4 entity maintaining a coordinated quality improvement program under this 5 section if the termination was on the basis of quality of care б concerns, introduction into evidence of information created, collected, 7 or maintained by the quality improvement committees of the subject 8 entity, which may be under terms of a protective order as specified by 9 10 the court; (e) in any civil action, disclosure of the fact that staff privileges were terminated or restricted, including the specific 11 12 restrictions imposed, if any and the reasons for the restrictions; or 13 (f) in any civil action, discovery and introduction into evidence of 14 the patient's medical records required by rule of the department of 15 health to be made regarding the care and treatment received.

16 (5) Information and documents created specifically for, and 17 collected and maintained by, a quality improvement committee are exempt 18 from disclosure under chapter 42.56 RCW.

(6) A coordinated quality improvement program may share information 19 and documents, including complaints and incident reports, created 20 21 specifically for, and collected and maintained by, a quality 22 improvement committee or a peer review committee under RCW 4.24.250 with one or more other coordinated quality improvement programs 23 24 maintained in accordance with this section or with RCW 70.41.200, a coordinated quality improvement committee maintained by an ambulatory 25 26 surgical facility under section 8 of this act, a quality assurance 27 committee maintained in accordance with RCW 18.20.390 or 74.42.640, or a peer review committee under RCW 4.24.250, for the improvement of the 28 quality of health care services rendered to patients and the 29 identification and prevention of medical malpractice. The privacy 30 protections of chapter 70.02 RCW and the federal health insurance 31 portability and accountability act of 1996 and its implementing 32 regulations apply to the sharing of individually identifiable patient 33 information held by a coordinated quality improvement program. Any 34 rules necessary to implement this section shall meet the requirements 35 of applicable federal and state privacy laws. 36 Information and 37 documents disclosed by one coordinated quality improvement program to another coordinated quality improvement program or a peer review 38

committee under RCW 4.24.250 and any information and documents created or maintained as a result of the sharing of information and documents shall not be subject to the discovery process and confidentiality shall be respected as required by subsection (4) of this section and RCW 4.24.250.

6 (7) The department of health shall adopt rules as are necessary to 7 implement this section.

8 Sec. 22. RCW 70.41.200 and 2005 c 291 s 3 and 2005 c 33 s 7 are 9 each reenacted and amended to read as follows:

10 (1) Every hospital shall maintain a coordinated quality improvement 11 program for the improvement of the quality of health care services 12 rendered to patients and the identification and prevention of medical 13 malpractice. The program shall include at least the following:

(a) The establishment of a quality improvement committee with the 14 responsibility to review the services rendered in the hospital, both 15 16 retrospectively and prospectively, in order to improve the quality of 17 medical care of patients and to prevent medical malpractice. The committee shall oversee and coordinate the quality improvement and 18 19 medical malpractice prevention program and shall ensure that 20 information gathered pursuant to the program is used to review and to 21 revise hospital policies and procedures;

(b) A medical staff privileges sanction procedure through which credentials, physical and mental capacity, and competence in delivering health care services are periodically reviewed as part of an evaluation of staff privileges;

(c) The periodic review of the credentials, physical and mental capacity, and competence in delivering health care services of all persons who are employed or associated with the hospital;

(d) A procedure for the prompt resolution of grievances by patients
 or their representatives related to accidents, injuries, treatment, and
 other events that may result in claims of medical malpractice;

(e) The maintenance and continuous collection of information
 concerning the hospital's experience with negative health care outcomes
 and incidents injurious to patients, patient grievances, professional
 liability premiums, settlements, awards, costs incurred by the hospital
 for patient injury prevention, and safety improvement activities;

1 (f) The maintenance of relevant and appropriate information 2 gathered pursuant to (a) through (e) of this subsection concerning 3 individual physicians within the physician's personnel or credential 4 file maintained by the hospital;

5 (g) Education programs dealing with quality improvement, patient 6 safety, medication errors, injury prevention, staff responsibility to 7 report professional misconduct, the legal aspects of patient care, 8 improved communication with patients, and causes of malpractice claims 9 for staff personnel engaged in patient care activities; and

10 (h) Policies to ensure compliance with the reporting requirements 11 of this section.

12 (2) Any person who, in substantial good faith, provides information 13 to further the purposes of the quality improvement and medical 14 malpractice prevention program or who, in substantial good faith, participates on the quality improvement committee shall not be subject 15 to an action for civil damages or other relief as a result of such 16 17 activity. Any person or entity participating in a coordinated quality improvement program that, in substantial good faith, shares information 18 or documents with one or more other programs, committees, or boards 19 under subsection (8) of this section is not subject to an action for 20 21 civil damages or other relief as a result of the activity. For the 22 purposes of this section, sharing information is presumed to be in substantial good faith. However, the presumption may be rebutted upon 23 24 showing of clear, cogent, and convincing evidence that а the 25 information shared was knowingly false or deliberately misleading.

(3) Information and documents, including complaints and incident 26 27 reports, created specifically for, and collected and maintained by, a quality improvement committee are not subject to review or disclosure, 28 except as provided in this section, or discovery or introduction into 29 evidence in any civil action, and no person who was in attendance at a 30 31 meeting of such committee or who participated in the creation, 32 collection, or maintenance of information or documents specifically for the committee shall be permitted or required to testify in any civil 33 34 action as to the content of such proceedings or the documents and information prepared specifically for the committee. This subsection 35 does not preclude: (a) In any civil action, the discovery of the 36 37 identity of persons involved in the medical care that is the basis of 38 the civil action whose involvement was independent of any quality

improvement activity; (b) in any civil action, the testimony of any 1 2 person concerning the facts which form the basis for the institution of such proceedings of which the person had personal knowledge acquired 3 independently of such proceedings; (c) in any civil action by a health 4 care provider regarding the restriction or revocation of that 5 individual's clinical or staff privileges, introduction into evidence б 7 information collected and maintained by quality improvement committees regarding such health care provider; (d) in any civil action, 8 disclosure of the fact that staff privileges were terminated or 9 restricted, including the specific restrictions imposed, if any and the 10 reasons for the restrictions; or (e) in any civil action, discovery and 11 introduction into evidence of the patient's medical records required by 12 13 regulation of the department of health to be made regarding the care 14 and treatment received.

15 (4) Each quality improvement committee shall, on at least a 16 semiannual basis, report to the governing board of the hospital in 17 which the committee is located. The report shall review the quality 18 improvement activities conducted by the committee, and any actions 19 taken as a result of those activities.

(5) The department of health shall adopt such rules as are deemedappropriate to effectuate the purposes of this section.

22 (6) The medical quality assurance commission or the board of osteopathic medicine and surgery, as appropriate, may review and audit 23 24 the records of committee decisions in which a physician's privileges 25 are terminated or restricted. Each hospital shall produce and make accessible to the commission or board the appropriate records and 26 27 otherwise facilitate the review and audit. Information so gained shall not be subject to the discovery process and confidentiality shall be 28 respected as required by subsection (3) of this section. Failure of a 29 hospital to comply with this subsection is punishable by a civil 30 penalty not to exceed two hundred fifty dollars. 31

(7) The department, the joint commission on accreditation of health care organizations, and any other accrediting organization may review and audit the records of a quality improvement committee or peer review committee in connection with their inspection and review of hospitals. Information so obtained shall not be subject to the discovery process, and confidentiality shall be respected as required by subsection (3) of 1 this section. Each hospital shall produce and make accessible to the 2 department the appropriate records and otherwise facilitate the review 3 and audit.

(8) A coordinated quality improvement program may share information 4 5 and documents, including complaints and incident reports, created specifically for, and collected and maintained by, a quality 6 7 improvement committee or a peer review committee under RCW 4.24.250 8 with one or more other coordinated quality improvement programs maintained in accordance with this section or RCW 43.70.510, a 9 coordinated quality improvement committee maintained by an ambulatory 10 surgical facility under section 8 of this act, a quality assurance 11 12 committee maintained in accordance with RCW 18.20.390 or 74.42.640, or 13 a peer review committee under RCW 4.24.250, for the improvement of the 14 quality of health care services rendered to patients and the identification and prevention of medical malpractice. The privacy 15 protections of chapter 70.02 RCW and the federal health insurance 16 portability and accountability act of 1996 and its implementing 17 regulations apply to the sharing of individually identifiable patient 18 information held by a coordinated quality improvement program. 19 Any rules necessary to implement this section shall meet the requirements 20 21 of applicable federal and state privacy laws. Information and 22 documents disclosed by one coordinated quality improvement program to another coordinated quality improvement program or a peer review 23 24 committee under RCW 4.24.250 and any information and documents created 25 or maintained as a result of the sharing of information and documents shall not be subject to the discovery process and confidentiality shall 26 27 be respected as required by subsection (3) of this section, RCW 18.20.390 (6) and (8), 74.42.640 (7) and (9), and 4.24.250. 28

(9) A hospital that operates a nursing home as defined in RCW 18.51.010 may conduct quality improvement activities for both the hospital and the nursing home through a quality improvement committee under this section, and such activities shall be subject to the provisions of subsections (2) through (8) of this section.

34 (10) Violation of this section shall not be considered negligence 35 per se.

36 **Sec. 23.** RCW 18.130.070 and 2006 c 99 s 2 are each amended to read 37 as follows:

(1)(a) The secretary shall adopt rules requiring every license 1 2 holder to report to the appropriate disciplining authority any conviction, determination, or finding that another license holder has 3 committed an act which constitutes unprofessional conduct, or to report 4 information to the disciplining authority, an impaired practitioner 5 program, or voluntary substance abuse monitoring program approved by 6 7 the disciplining authority, which indicates that the other license holder may not be able to practice his or her profession with 8 reasonable skill and safety to consumers as a result of a mental or 9 10 physical condition.

(b) The secretary may adopt rules to require other persons, including corporations, organizations, health care facilities, impaired practitioner programs, or voluntary substance abuse monitoring programs approved by a disciplining authority, and state or local government agencies to report:

(i) Any conviction, determination, or finding that a license holderhas committed an act which constitutes unprofessional conduct; or

(ii) Information to the disciplining authority, an impaired practitioner program, or voluntary substance abuse monitoring program approved by the disciplining authority, which indicates that the license holder may not be able to practice his or her profession with reasonable skill and safety to consumers as a result of a mental or physical condition.

24 (c) If a report has been made by a hospital to the department pursuant to RCW 70.41.210 or by an ambulatory surgical facility 25 26 pursuant to section 12 of this act, a report to the disciplining 27 authority is not required. To facilitate meeting the intent of this section, the cooperation of agencies of the federal government is 28 requested by reporting any conviction, determination, or finding that 29 a federal employee or contractor regulated by the disciplining 30 authorities enumerated in this chapter has committed an act which 31 32 constituted unprofessional conduct and reporting any information which indicates that a federal employee or contractor regulated by the 33 disciplining authorities enumerated in this chapter may not be able to 34 35 practice his or her profession with reasonable skill and safety as a 36 result of a mental or physical condition.

37 (d) Reporting under this section is not required by:

1 (i) Any entity with a peer review committee, quality improvement 2 committee or other similarly designated professional review committee, 3 or by a license holder who is a member of such committee, during the 4 investigative phase of the respective committee's operations if the 5 investigation is completed in a timely manner; or

6 (ii) An impaired practitioner program or voluntary substance abuse 7 monitoring program approved by a disciplining authority under RCW 8 18.130.175 if the license holder is currently enrolled in the treatment 9 program, so long as the license holder actively participates in the 10 treatment program and the license holder's impairment does not 11 constitute a clear and present danger to the public health, safety, or 12 welfare.

13 (2) If a person fails to furnish a required report, the 14 disciplining authority may petition the superior court of the county in 15 which the person resides or is found, and the court shall issue to the 16 person an order to furnish the required report. A failure to obey the 17 order is a contempt of court as provided in chapter 7.21 RCW.

(3) A person is immune from civil liability, whether direct or
derivative, for providing information to the disciplining authority
pursuant to the rules adopted under subsection (1) of this section.

21 (4)(a) The holder of a license subject to the jurisdiction of this 22 chapter shall report to the disciplining authority:

(i) Any conviction, determination, or finding that he or she has committed unprofessional conduct or is unable to practice with reasonable skill or safety; and

26 (ii) Any disqualification from participation in the federal 27 medicare program, under Title XVIII of the federal social security act 28 or the federal medicaid program, under Title XIX of the federal social 29 security act.

30 (b) Failure to report within thirty days of notice of the 31 conviction, determination, finding, or disqualification constitutes 32 grounds for disciplinary action.

33 **Sec. 24.** RCW 18.71.0195 and 2005 c 274 s 227 are each amended to 34 read as follows:

(1) The contents of any report filed under RCW 18.130.070 shall be
 confidential and exempt from public disclosure pursuant to chapter
 42.56 RCW, except that it may be reviewed (a) by the licensee involved

or his or her counsel or authorized representative who may submit any additional exculpatory or explanatory statements or other information, which statements or other information shall be included in the file, or (b) by a representative of the commission, or investigator thereof, who has been assigned to review the activities of a licensed physician.

6 Upon a determination that a report is without merit, the 7 commission's records may be purged of information relating to the 8 report.

9 (2) Every individual, medical association, medical society, hospital, ambulatory surgical facility, medical service bureau, health 10 insurance carrier or agent, professional liability insurance carrier, 11 professional standards review organization, agency of the federal, 12 13 state, or local government, or the entity established by RCW 18.71.300 and its officers, agents, and employees are immune from civil 14 liability, whether direct or derivative, for providing information to 15 16 the commission under RCW 18.130.070, or for which an individual health 17 care provider has immunity under the provisions of RCW 4.24.240, 4.24.250, or 4.24.260. 18

19 Sec. 25. RCW 42.56.360 and 2006 c 209 s 9 and 2006 c 8 s 112 are 20 each reenacted and amended to read as follows:

(1) The following health care information is exempt from disclosureunder this chapter:

(a) Information obtained by the board of pharmacy as provided inRCW 69.45.090;

(b) Information obtained by the board of pharmacy or the department
of health and its representatives as provided in RCW 69.41.044,
69.41.280, and 18.64.420;

(c) Information and documents created specifically for, and 28 collected and maintained by a quality improvement committee under RCW 29 30 43.70.510, section 9 of this act, or 70.41.200, or by a peer review committee under RCW 4.24.250, or by a quality assurance committee 31 pursuant to RCW 74.42.640 or 18.20.390, and notifications or reports of 32 adverse events or incidents made under RCW 70.56.020 or 70.56.040, 33 regardless of which agency is in possession of the information and 34 35 documents;

36 (d)(i) Proprietary financial and commercial information that the 37 submitting entity, with review by the department of health,

specifically identifies at the time it is submitted and that is provided to or obtained by the department of health in connection with an application for, or the supervision of, an antitrust exemption sought by the submitting entity under RCW 43.72.310;

5 (ii) If a request for such information is received, the submitting 6 entity must be notified of the request. Within ten business days of 7 receipt of the notice, the submitting entity shall provide a written 8 statement of the continuing need for confidentiality, which shall be 9 provided to the requester. Upon receipt of such notice, the department 10 of health shall continue to treat information designated under this 11 subsection (1)(d) as exempt from disclosure;

(iii) If the requester initiates an action to compel disclosure under this chapter, the submitting entity must be joined as a party to demonstrate the continuing need for confidentiality;

(e) Records of the entity obtained in an action under RCW 18.71.300 through 18.71.340;

(f) Except for published statistical compilations and reports relating to the infant mortality review studies that do not identify individual cases and sources of information, any records or documents obtained, prepared, or maintained by the local health department for the purposes of an infant mortality review conducted by the department of health under RCW 70.05.170; and

(g) Complaints filed under chapter 18.130 RCW after July 27, 1997,
to the extent provided in RCW 18.130.095(1).

(2) Chapter 70.02 RCW applies to public inspection and copying ofhealth care information of patients.

27 Sec. 26. RCW 18.71.017 and 2000 c 171 s 23 are each amended to 28 read as follows:

(1) The commission may adopt such rules as are not inconsistent 29 30 with the laws of this state as may be determined necessary or proper to 31 carry out the purposes of this chapter. The commission is the successor in interest of the board of medical examiners and the medical 32 disciplinary board. All contracts, undertakings, agreements, rules, 33 regulations, and policies continue in full force and effect on July 1, 34 35 1994, unless otherwise repealed or rejected by this chapter or by the 36 commission.

(2) The commission may adopt rules governing the administration of
 sedation and anesthesia in the offices of persons licensed under this
 chapter, including necessary training and equipment.

4 **Sec. 27.** RCW 18.57.005 and 1986 c 259 s 94 are each amended to 5 read as follows:

6

The board shall have the following powers and duties:

7 (1) To administer examinations to applicants for licensure under8 this chapter;

9 (2) To make such rules and regulations as are not inconsistent with 10 the laws of this state as may be deemed necessary or proper to carry 11 out the purposes of this chapter;

12 To establish and administer requirements for continuing (3) professional education as may be necessary or proper to insure the 13 public health and safety as a prerequisite to granting and renewing 14 licenses under this chapter: PROVIDED, That such rules shall not 15 16 require a licensee under this chapter to engage in continuing education related to or provided by any specific branch, school, or philosophy of 17 medical practice or its political and/or professional organizations, 18 19 associations, or societies;

20 (4) <u>To adopt rules governing the administration of sedation and</u> 21 <u>anesthesia in the offices of persons licensed under this chapter,</u> 22 <u>including necessary training and equipment;</u>

23 (5) To keep an official record of all its proceedings, which record 24 shall be evidence of all proceedings of the board which are set forth 25 therein.

26 **Sec. 28.** RCW 18.22.015 and 1990 c 147 s 5 are each amended to read 27 as follows:

28 The board shall:

29 (1) Administer all laws placed under its jurisdiction;

30 (2) Prepare, grade, and administer or determine the nature,
 31 grading, and administration of examinations for applicants for
 32 podiatric physician and surgeon licenses;

33 (3) Examine and investigate all applicants for podiatric physician 34 and surgeon licenses and certify to the secretary all applicants it 35 judges to be properly qualified; (4) Adopt any rules which it considers necessary or proper to carry
 out the purposes of this chapter;

3 (5) Adopt rules governing the administration of sedation and 4 anesthesia in the offices of persons licensed under this chapter, 5 including necessary training and equipment;

6 (6) Determine which schools of podiatric medicine and surgery will 7 be approved.

8 <u>NEW SECTION.</u> Sec. 29. Except for section 7 of this act, this act 9 takes effect July 1, 2009.

10 <u>NEW SECTION.</u> **Sec. 30.** The secretary of health may take the 11 necessary steps to ensure that this act is implemented on its effective 12 date.

13 <u>NEW SECTION.</u> Sec. 31. Sections 1 through 6 and 8 through 19 of 14 this act constitute a new chapter in Title 70 RCW.

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