Title 69  
FOOD, DRUGS, COSMETICS, AND POISONS

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Chapter 69.04 RCW
INTRASTATE COMMERCE IN FOOD, DRUGS, AND COSMETICS

(Formerly: Food, drug, and cosmetic act)

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2012
69.04.001 Statement of purpose. This chapter is intended to enact state legislation (1) which safeguards the public health and promotes the public welfare by protecting the consuming public from (a) potential injury by product use; (b) products that are adulterated; or (c) products that have been produced under unsanitary conditions, and the purchasing public from injury by merchandising deceit flowing from intrastate commerce in food, drugs, devices, and cosmetics; and (2) which is uniform, as provided in this chapter, with the federal food, drug, and cosmetic act; and with the federal trade commission act, to the extent it expressly outlaws the false advertisement of food, drugs, devices, and cosmetics; and (3) which thus promotes uniformity of such law and its administration and enforcement, in and throughout the United States. [1991 c 162 § 1; 1945 c 257 § 2; Rem. Supp. 1945 § 6163-51.]

Conformity with federal regulations: RCW 69.04.190 and 69.04.200.

69.04.002 Introductory. For the purposes of this chapter, terms shall apply as herein defined unless the context clearly indicates otherwise. [1945 c 257 § 3; Rem. Supp. 1945 § 6163-52.]


69.04.004 "Intrastate commerce." The term "intrastate commerce" means any and all commerce within the state of Washington and subject to the jurisdiction thereof; and includes the operation of any business or service establishment. [1945 c 257 § 5; Rem. Supp. 1945 § 6163-54.]

69.04.005 "Sale." The term "sale" means any and every sale and includes (1) manufacture, processing, packing, canning, bottling, or any other production, preparation, or putting up; (2) exposure, offer, or any other proffer; (3) holding, storing, or any other possessing; (4) dispensing, giving, delivering, serving, or any other supplying; and (5) applying, administering, or any other using. [1945 c 257 § 6; Rem. Supp. 1945 § 6163-55.]
69.04.006 "Director." The term "director" means the director of the department of agriculture of the state of Washington and his or her duly authorized representatives. [2012 c 117 § 328; 1945 c 257 § 7; Rem. Supp. 1945 § 6163-56.]

Director of agriculture, general duties: Chapter 43.23 RCW.

69.04.007 "Person." The term "person" includes individual, partnership, corporation, and association. [1945 c 257 § 8; Rem. Supp. 1945 § 6163-57.]

69.04.008 "Food." The term "food" means (1) articles used for food or drink for people or other animals, (2) bottled water, (3) chewing gum, and (4) articles used for components of any such article. [1992 c 34 § 2; 1945 c 257 § 9; Rem. Supp. 1945 § 6163-58.]

Additional notes found at www.leg.wa.gov

69.04.009 "Drugs." The term "drug" means (1) articles recognized in the official United States pharmacopoeia, official homeopathic pharmacopoeia of the United States, or official national formulary, or any supplement to any of them; and (2) articles intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in human beings or other animals; and (3) articles (other than food) intended to affect the structure or any function of components of human beings or other animals; and (4) articles intended for use as a component of any article specified in clause (1), (2), or (3); but does not include devices or their components, parts, or accessories. [2009 c 549 § 1018; 1945 c 257 § 10; Rem. Supp. 1945 § 6163-59. Prior: 1907 c 211 § 2.]

69.04.010 "Device." The term "device" (except when used in RCW 69.04.016 and in RCW 69.04.040(10), 69.04.270, 69.04.690, and in RCW 69.04.470 as used in the sentence "(as compared with other words, statements, designs, or devices, in the labeling") means instruments, apparatus, and contrivances, including their components, parts and accessories, intended (1) for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in human beings or other animals; or (2) to affect the structure or any function of the body of human beings or other animals. [2009 c 549 § 1019; 1945 c 257 § 11; Rem. Supp. 1945 § 6163-60.]

69.04.011 "Cosmetic." The term "cosmetic" means (1) articles intended to be rubbed, poured, sprinkled, or sprayed on, introduced into, or otherwise applied to the human body or any part thereof for cleansing, beautifying, promoting attractiveness, or altering the appearance, and (2) articles intended for use as a component of any such article; except that such term shall not include soap. [1945 c 257 § 12; Rem. Supp. 1945 § 6163-61.]

69.04.012 "Official compendium." The term "official compendium" mean the official United States pharmacopoeia, official homeopathic pharmacopoeia of the United States, official national formulary, or any supplement to any of them. [1945 c 257 § 13; Rem. Supp. 1945 § 6163-62.]

69.04.013 "Label." The term "label" means a display of written, printed, or graphic matter upon the immediate container of any article; and a requirement made by or under authority of this chapter that any word, statement, or other information appears on the label shall not be considered to be complied with unless such word, statement, or other information also appears on the outside container or wrapper, if any there be, of the retail package of such article, or is easily legible through the outside container or wrapper. [1945 c 257 § 14; Rem. Supp. 1945 § 6163-63.]

69.04.014 "Immediate container." The term "immediate container" does not include package liners. [1945 c 257 § 15; Rem. Supp. 1945 § 6163-64.]

69.04.015 "Labeling." The term "labeling" means all labels and other written, printed, or graphic matter (1) upon any article or any of its containers or wrappers, or (2) accompanying such article. [1945 c 257 § 16; Rem. Supp. 1945 § 6163-65.]

Crimes relating to labeling: Chapter 9.16 RCW, RCW 69.40.055.

69.04.016 "Misleading labeling or advertisement," how determined. If any article is alleged to be misbranded because the labeling is misleading, or if an advertisement is alleged to be false because it is misleading, then in determining whether the labeling or advertisement is misleading there shall be taken into account (among other things) not only representations made or suggested by statement, word, design, device, sound, or any combination thereof, but also the extent to which the labeling or advertisement fails to reveal facts material in the light of such representations or material with respect to consequences which may result from the use of the article to which the labeling or advertisement relates under the conditions of use prescribed in the labeling or advertisement thereof or under such conditions of use as are customary or usual. [1945 c 257 § 17; Rem. Supp. 1945 § 6163-66.]

Crimes relating to advertising: Chapter 9.04 RCW.

69.04.017 "Antiseptic" as germicide. The representation of a drug, in its labeling or advertisement, as an antiseptic shall be considered to be a representation that it is a germicide, except in the case of a drug purporting to be, or represented as, an antiseptic for inhibitory use as a wet dressing, ointment, dusting powder, or such other use as involves prolonged contact with the body. [1945 c 257 § 18; Rem. Supp. 1945 § 6163-67.]

69.04.018 "New drug" defined. The term "new drug" means (1) any drug the composition of which is such that such drug is not generally recognized, among experts qualified by scientific training and experience to evaluate the safety of drugs, as safe for use under the conditions prescribed, recommended, or suggested in the labeling thereof; or (2) any drug the composition of which is such that such drug, as a result of investigations to determine its safety for use under such conditions, has become so recognized, but which has not, otherwise than in such investigations, been used to a material extent or for a material time under such conditions: PROVIDED, That no drug in use on the effective date of this chapter shall be regarded as a new drug. [1945 c 257 § 19; Rem. Supp. 1945 § 6163-68.]

Additional notes found at www.leg.wa.gov
69.04.019 "Advertisement." The term "advertisement" means all representations, other than by labeling, for the purpose of inducing, or which are likely to induce, directly or indirectly the purchase of food, drugs, devices, or cosmetics. [1945 c 257 § 20; Rem. Supp. 1945 § 6163-69.]

69.04.020 "Contaminated with filth." The term "contaminated with filth" applies to any food, drug, device, or cosmetic not securely protected from dust, dirt, and as far as may be necessary by all reasonable means, from all foreign or injurious contaminations. [1945 c 257 § 21; Rem. Supp. 1945 § 6163-70.]

69.04.021 "Package." The word "package" shall include, and be construed to include, wrapped meats enclosed in papers or other materials as prepared by the manufacturers thereof for sale. [1963 c 198 § 8.]

69.04.022 "Pesticide chemical." The term "pesticide chemical" means any substance defined as an economic poison and/or agricultural pesticide in Title 15 RCW as now enacted or hereafter amended. [1963 c 198 § 9.]

69.04.023 "Raw agricultural commodity." The term "raw agricultural commodity" means any food in its raw or natural state, including all fruits that are washed, colored or otherwise treated in their unpeeled natural form prior to marketing. [1963 c 198 § 10.]

69.04.024 "Food additive," "safe." (1) The term "food additive" means any substance the intended use of which results or may reasonably be expected to result, directly or indirectly, in its becoming a component or otherwise affecting the characteristics of any food (including any substance intended for use in producing, manufacturing, packing, processing, preparing, treating, packaging, transporting, or holding food; and including any source of radiation intended for any such use), if such substance generally is recognized, among experts qualified by scientific training and experience to evaluate its safety, as having been adequately shown through scientific procedures (or, in the case of a substance used in food prior to January 1, 1958; through either scientific procedures or experience based on common use in food) to be unsafe under the conditions of its intended use; except that such term does not include; (a) a pesticide chemical in or on a raw agricultural commodity; or (b) a pesticide chemical to the extent that it is intended for use or is used in the production, storage, or transportation of any raw agricultural commodity; or (c) a color additive.

(2) The term "safe" as used in the food additive definition has reference to the health of human beings or animals. [2009 c 549 § 1020; 1963 c 198 § 11.]

69.04.025 "Color additive," "color." (1) The term "color additive" means a material which (a) is a dye, pigment, or other substance made by a process of synthesis or similar artifice, or extracted, isolated, or otherwise derived, with or without intermediate or final change of identity, from a vegetable, animal, mineral, or other source, and (b) when added or applied to a food is capable (alone or through reaction with other substance) of imparting color thereto; except that such term does not include any material which the director, by regulation, determines is used (or intended to be used) solely for a purpose or purposes other than coloring.

(2) The term "color" includes black, white, and intermediate grays.

(3) Nothing in subsection (1) hereof shall be construed to apply to any pesticide chemical, soil or plant nutrient, or other agricultural chemical solely because of its effect in aiding, retarding, or otherwise affecting, directly or indirectly, the growth or other natural physiological processes of produce of the soil and thereby affecting its color, whether before or after harvest. [1963 c 198 § 12.]

69.04.040 Prohibited acts. The following acts and the causing thereof are hereby prohibited:

(1) The sale in intrastate commerce of any food, drug, device, or cosmetic that is adulterated or misbranded.

(2) The adulteration or misbranding of any food, drug, device, or cosmetic in intrastate commerce.

(3) The receipt in intrastate commerce of any food, drug, device, or cosmetic that is adulterated or misbranded, and the sale thereof in such commerce for pay or otherwise.

(4) The introduction or delivery for introduction into intrastate commerce of (a) any food in violation of RCW 69.04.350; or (b) any new drug in violation of RCW 69.04.570.

(5) The dissemination within this state, in any manner or by any means or through any medium, of any false advertisement.

(6) The refusal to permit (a) entry and the taking of a sample or specimen or the making of any investigation or examination as authorized by RCW 69.04.780; or (b) access to or copying of any record as authorized by RCW 69.04.810.

(7) The refusal to permit entry or inspection as authorized by RCW 69.04.820.

(8) The removal, mutilation, or violation of an embargo notice as authorized by RCW 69.04.110.

(9) The giving of a guaranty or undertaking in intrastate commerce, referred to in RCW 69.04.080, that is false.

(10) The forging, counterfeiting, simulating, or falsely representing, or without proper authority, using any mark, stamp, tag, label, or other identification device authorized or required by regulations promulgated under RCW 69.04.350.

(11) The alteration, mutilation, destruction, obliteration, or removal of the whole or any part of the labeling of a food, drug, device, or cosmetic, or the doing of any other act with respect to a food, drug, device, or cosmetic, or the labeling or advertisement thereof, which results in a violation of this chapter.

(12) The using in intrastate commerce, in the labeling or advertisement of any drug, of any representation or suggestion that an application with respect to such drug is effective under section 505 of the federal act or under RCW 69.04.570, or that such drug complies with the provisions of either such section. [1945 c 257 § 22; Rem. Supp. 1945 § 6163-71. Prior: 1917 c 168 § 1; 1907 c 211 § 1; 1901 c 94 § 1.]

69.04.050 Remedy by injunction. (1) In addition to the remedies hereinafter provided the director is hereby authorized to apply to the superior court of Thurston county for, and such court shall have jurisdiction upon prompt hearing
and for cause shown to grant, a temporary or permanent injunction restraining any person from violating any provision of RCW 69.04.040; without proof that an adequate remedy at law does not exist.

(2) Whenever it appears to the satisfaction of the court in the case of a newspaper, magazine, periodical, or other publication, published at regular intervals (a) that restraining the dissemination of a false advertisement in any particular issue of such publication would delay the delivery of such issue after the regular time therefor, and (b) that such delay would be due to the method by which the manufacture and distribution of such publication is customarily conducted by the publisher in accordance with sound business practice, and not to any method or device adopted for the evasion of this section or to prevent or delay the issuance of an injunction or restraining order with respect to such false advertisement or any other advertisement, the court shall exclude such issue from the operation of the restraining order or injunction. [1945 c 257 § 23; Rem. Supp. 1945 § 6163-72.]

Injunctions, generally: Chapter 7.40 RCW.

69.04.060 Criminal penalty for violations. Any person who violates any provision of RCW 69.04.040 is guilty of a misdemeanor and shall on conviction thereof be subject to the following penalties:

(1) A fine of not more than two hundred dollars; or

(2) If the violation is committed after a conviction of such person under this section has become final, imprisonment for not more than thirty days, or a fine of not more than five hundred dollars, or both such imprisonment and fine. [2003 c 53 § 314; 1945 c 257 § 24; Rem. Supp. 1945 § 6163-73. Prior: 1907 c 211 § 12; 1901 c 94 § 11.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.04.070 Additional penalty. Notwithstanding the provisions of RCW 69.04.060, a person who violates RCW 69.04.040 with intent to defraud or mislead is guilty of a misdemeanor and the penalty shall be imprisonment for not more than ninety days, or a fine of not more than one thousand dollars, or both such imprisonment and fine. [2003 c 53 § 315; 1945 c 257 § 25; Rem. Supp. 1945 § 6163-74.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.04.080 Avoidance of penalty. No person shall be subject to the penalties of RCW 69.04.060:

(1) For having violated RCW 69.04.040(3), if he or she establishes that he or she received and sold such article in good faith, unless he or she refuses on request of the director to furnish the name and address of the person in the state of Washington from whom he or she received such article and copies of all available documents pertaining to his or her receipt thereof; or

(2) For having violated RCW 69.04.040 (1), (3), or (4), if he or she establishes a guaranty or undertaking signed by, and containing the name and address of, the person in the state of Washington from whom he or she received such article in good faith, to the effect that such article complies with this chapter; or

(3) For having violated RCW 69.04.040(5), if he or she establishes a guaranty or undertaking signed by, and containing the name and address of, the person in the state of Washington from whom he or she received such advertisement in good faith, to the effect that such advertisement complies with this chapter; or

(4) For having violated RCW 69.04.040(9), if he or she establishes that he or she gave such guaranty or undertaking in good faith and in reliance on a guaranty or undertaking to him or her, which guaranty or undertaking was to the same effect and was signed by, and contained the name and address of, a person in the state of Washington. [2012 c 117 § 329; 1945 c 257 § 26; Rem. Supp. 1945 § 6163-75.]

69.04.090 Liability of disseminator of advertisement. No publisher, radio broadcast licensee, advertising agency, or agency or medium for the dissemination of an advertisement, except the manufacturer, packer, distributor, or seller of the article to which the advertisement relates, shall be subject to the penalties of RCW 69.04.060 by reason of his or her dissemination of any false advertisement, unless he or she has refused on the request of the director to furnish the name and address of the manufacturer, packer, distributor, seller, or advertising agency in the state of Washington, who caused him or her to disseminate such false advertisement. [2012 c 117 § 330; 1945 c 257 § 27; Rem. Supp. 1945 § 6163-76.]

69.04.100 Condemnation of adulterated or misbranded article. Whenever the director shall find in intrastate commerce an article subject to this chapter which is so adulterated or misbranded that it is unfit or unsafe for human use and its immediate condemnation is required to protect the public health, such article is hereby declared to be a nuisance and the director is hereby authorized forthwith to destroy such article or to render it unsalable for human use. [1945 c 257 § 28; Rem. Supp. 1945 § 6163-77.]

69.04.110 Embargo of articles. Whenever the director shall find, or shall have probable cause to believe, that an article subject to this chapter is in intrastate commerce in violation of this chapter, and that its embargo under this section is required to protect the consuming or purchasing public, due to its being adulterated or misbranded, or to otherwise protect the public from injury, or possible injury, he or she is hereby authorized to affix to such article a notice of its embargo and against its sale in intrastate commerce, without permission given under this chapter. But if, after such article has been so embargoed, the director shall find that such article does not involve a violation of this chapter, such embargo shall be forthwith removed. [1991 c 162 § 3; 1975 1st ex.s. c 7 § 25; 1945 c 257 § 29; Rem. Supp. 1945 § 6163-78.]

Purpose of section: See RCW 69.04.398.

69.04.120 Procedure on embargo. When the director has embargoed an article, he or she shall, forthwith and without delay and in no event later than thirty days after the affixing of notice of its embargo, petition the superior court for an order affirming the embargo. The court then has jurisdiction, for cause shown and after prompt hearing to any claimant of the embargoed article, to issue an order which directs the removal of the embargo or the destruction or the correction of
and release of the article. An order for destruction or correction and release shall contain such provision for the payment of pertinent court costs and fees and administrative expenses as is equitable and which the court deems appropriate in the circumstances. An order for correction and release may contain such provision for a bond as the court finds indicated in the circumstances. [1991 c 162 § 4; 1983 c 95 § 8; 1945 c 257 § 30; Rem. Supp. 1945 § 6163-79.]

**69.04.123 Exception to petition requirement under RCW 69.04.120.** The director need not petition the superior court as provided for in RCW 69.04.120 if the owner or claimant of such food or food products agrees in writing to the disposition of such food or food products as the director may order. [1995 c 374 § 20.]

Additional notes found at www.leg.wa.gov

**69.04.130 Petitions may be consolidated.** Two or more petitions under RCW 69.04.120, which pend at the same time and which present the same issue and claimant hereunder, shall be consolidated for simultaneous determination by one court of jurisdiction, upon application to any court of jurisdiction by the director or by such claimant. [1945 c 257 § 31; Rem. Supp. 1945 § 6163-80.]

**69.04.140 Claimant entitled to sample.** The claimant in any proceeding by petition under RCW 69.04.120 shall be entitled to receive a representative sample of the article subject to such proceeding, upon application to the court of jurisdiction made at any time after such petition and prior to the hearing thereon. [1945 c 257 § 32; Rem. Supp. 1945 § 6163-81.]

**69.04.150 Damages not recoverable if probable cause existed.** No state court shall allow the recovery of damages from administrative action for condemnation under RCW 69.04.100 or for embargo under RCW 69.04.110, if the court finds that there was probable cause for such action. [1945 c 257 § 33; Rem. Supp. 1945 § 6163-82.]

**69.04.160 Prosecutions.** (1) It shall be the duty of each state attorney, county attorney, or city attorney to whom the director reports any violation of this chapter, or regulations promulgated under it, to cause appropriate proceedings to be instituted in the proper courts, without delay, and to be duly prosecuted as prescribed by law.

(2) Before any violation of this chapter is reported by the director to any such attorney for the institution of a criminal proceeding, the person against whom such proceeding is contemplated shall be given appropriate notice and an opportunity to present his or her views to the director, either orally or in writing, with regard to such contemplated proceeding. [2012 c 117 § 331; 1945 c 257 § 34; Rem. Supp. 1945 § 6163-83.]

**69.04.170 Minor infractions.** Nothing in this chapter shall be construed as requiring the director to report for the institution of proceedings under this chapter, minor violations of this chapter, whenever he or she believes that the public interest will be adequately served in the circumstances by a suitable written notice or warning. [2012 c 117 § 332; 1945 c 257 § 35; Rem. Supp. 1945 § 6163-84.]

**69.04.180 Proceedings to be in name of state.** All such proceedings for the enforcement, or to restrain violations, of this chapter shall be by and in the name of the state of Washington. [1945 c 257 § 36; Rem. Supp. 1945 § 6163-85.]

**69.04.190 Standards may be prescribed by regulations.** Whenever in the judgment of the director such action will promote honesty and fair dealing in the interest of consumers, he or she shall promulgate regulations fixing and establishing for any food, under its common or usual name so far as practicable, a reasonable definition and standard of identity, a reasonable standard of quality, and/or reasonable standards of fill of container. In prescribing any standard of fill of container, consideration shall be given to and due allowance shall be made for product or volume shrinkage or expansion unavoidable in good commercial practice, and need for packing and protective material. In prescribing any standard of quality for any canned fruit or canned vegetable, consideration shall be given to and due allowance shall be made for the differing characteristics of the several varieties thereof. In prescribing a definition and standard of identity for any food or class of food in which optional ingredients are permitted, the director shall, for the purpose of promoting honesty and fair dealing in the interest of consumers, designate the optional ingredients which shall be named on the label. [2012 c 117 § 333; 1945 c 257 § 37; Rem. Supp. 1945 § 6163-86. Prior: 1917 c 168 § 2.]

**69.04.200 Conformance with federal standards.** The definitions and standards of identity, the standards of quality and fill of container, and the label requirements prescribed by regulations promulgated under this section shall conform, insofar as practicable, with those prescribed by regulations promulgated under section 401 of the federal act and to the definitions and standards promulgated under the meat inspection act approved March 4, 1907, as amended. [1945 c 257 § 38; Rem. Supp. 1945 § 6163-87.]

*Revisor’s note: The language “this section” appears in 1945 c 257 § 38 but apparently refers to 1945 c 257 § 37 codified as RCW 69.04.190.

**69.04.205 Bacon—Packaging at retail to reveal quality and leanness.** All packaged bacon other than that packaged in cans shall be offered and exposed for sale and sold, within the state of Washington only at retail in packages which permit the buyer to readily view the quality and degree of leanness of the product. [1971 c 49 § 1.]

**69.04.206 Bacon—Rules, regulations, and standards—Withholding packaging use—Hearing—Final determination—Appeal.** The director of the department of agriculture is hereby authorized to promulgate rules, regulations, and standards for the implementation of RCW 69.04.205 through 69.04.207. If the director has reason to believe that any packaging method, package, or container in use or proposed for use with respect to the marketing of bacon is false or misleading in any particular, or does not meet the requirements of RCW 69.04.205, he or she may
direct that such use be withheld unless the packaging method, package, or container is modified in such manner as he deems necessary so that it will not be false or misleading. If the person, firm, or corporation using or proposing to use the packaging method, package, or container does not accept the determination of the director, he may prescribe so that it will not be false or misleading. If the use of the radiation was in conformity with a regulation or exemption in effect pursuant to RCW 69.04.394. [1963 c 198 § 1; 1945 c 257 § 39; Rem. Supp. 1945 § 6163-88. Prior: 1923 c 36 § 1; 1907 c 211 § 3; 1901 c 94 § 3.]

69.04.220 Food—Adulteration by abstraction, addition, substitution, etc. A food shall be deemed to be adulterated (1) if any valuable constituent has been in whole or in part omitted or abstracted therefrom; or (2) if any substance has been substituted wholly or in part therefor; or (3) if damage or inferiority has been concealed in any manner; or (4) if any substance has been added thereto or mixed or packed therewith so as to increase its bulk or weight, or reduce its quality or strength, or make it appear better or of greater value than it is. [1945 c 257 § 40; Rem. Supp. 1945 § 6163-89.]

69.04.231 Food—Adulteration by color additive. A food shall be deemed to be adulterated if it is, or it bears or contains a color additive which is unsafe within the meaning of RCW 69.04.396. [1963 c 198 § 5.]

69.04.240 Confectionery—Adulteration. A food shall be deemed to be adulterated if it is confectionery and it bears or contains any alcohol from natural or artificial alcohol flavoring in excess of one percent of the weight of the confection or any nonnutritive article or substance except harmless coloring, harmless flavoring, harmless resinous glaze not in excess of four-tenths of one percent, natural gum, and pectin. This section shall not apply to any chewing gum by reason of its containing harmless nonnutritive masticatory substances, or to any confection permitted to be sold by an endorsement from the liquor control board under RCW 66.24.360. [2007 c 226 § 3; 1984 c 78 § 2; 1945 c 257 § 42; Rem. Supp. 1945 § 6163-91. Prior: 1923 c 36 § 1, part; 1907 c 211 § 3, part.]


69.04.245 Poultry—Improper use of state’s geographic outline. Uncooked poultry is deemed to be misbranded if it is produced outside of this state but the label for the poultry contains the geographic outline of this state. [1989 c 257 § 2.]

Legislative findings—1989 c 257: "The legislature finds that: Poultry produced in this state is known throughout the state for its high quality; and one of the sources of that quality is the proximity of production centers to retail outlets in the state. The legislature also finds that labeling which misrepresents poultry produced elsewhere as being a product of this state may lead consumers to purchase products which they would not otherwise purchase. The legislature further finds that the presence of the geographic outline of this state on a label for poultry produced outside of the state misrepresents the product as having been produced in this state." [1989 c 257 § 1.]

69.04.250 Food—Misbranding by false label, etc. A food shall be deemed to be misbranded (1) if its labeling is false or misleading in any particular; or (2) if it is offered for sale under the name of another food; or (3) if it is an imitation of another food, unless its label bears, in type of uniform size and prominence, the word "imitation" and, immediately

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thereafter, the name of the food imitated; or (4) if its container is so made, formed or filled as to be misleading. [1945 c 257 § 43; Rem. Supp. 1945 § 6163-92. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.260 Packaged food—Misbranding. If a food is in package form, it shall be deemed to be misbranded, unless it bears a label containing (1) the name and place of business of the manufacturer, packer, or distributor; and (2) an accurate statement of the quantity of the contents in terms of weight, measure, or numerical count: PROVIDED, That under clause (2) of this section reasonable variations shall be permitted, and exemptions as to small packages shall be established, by regulations promulgated by the director. [1945 c 257 § 44; Rem. Supp. 1945 § 6163-93.]

69.04.270 Food—Misbranding by lack of prominent label. A food shall be deemed to be misbranded if any word, statement, or other information required by or under authority of this chapter to appear on the label or labeling is not prominently placed thereon with such conspicuousness (as compared with other words, statements, designs, or devices, in the labeling) and in such terms as to render it likely to be read and understood by the ordinary individual under customary conditions of purchase and use. [1945 c 257 § 45; Rem. Supp. 1945 § 6163-94.]

69.04.280 Food—Misbranding for nonconformity with standard of identity. If a food purports to be or is represented as a food for which a definition and standard of identity has been prescribed by regulations as provided by RCW 69.04.190, it shall be deemed to be misbranded unless (1) it conforms to such definition and standard, and (2) its label bears the name of the food specified in the definition and standard, and, insofar as may be required by such regulations, the common or usual name of each such ingredient; except that spices, flavorings, and colorings, other than those sold as such, may be designated as spices, flavorings, and colorings without naming each: PROVIDED, That, to the extent that compliance with the requirements of clause (2) of this section is impracticable, or results in deception or unfair competition, exemptions shall be established by regulations promulgated by the director. [1945 c 257 § 49; Rem. Supp. 1945 § 6163-98.]

69.04.315 Halibut—Misbranding by failure to show proper name. No person shall label or offer for sale any food fish product designated as halibut, with or without additional descriptive words unless such food fish product is Hippoglossus Hippoglossus or Hippoglossus Stenolepis. Any person violating the provisions of this section shall be guilty of misbranding under the provisions of this chapter. [1967 ex.s. c 79 § 1.]

69.04.320 Food—Misbranding by failure to show dietary properties. If a food purports to be or is represented for special dietary uses, it shall be deemed to be misbranded, unless its label bears such information concerning its vitamin, mineral and other dietary properties as is necessary in order to fully inform purchasers as to its value for such uses, as provided by regulations promulgated by the director, such regulations to conform insofar as practicable with regulations under section 403(j) of the federal act. [1945 c 257 § 50; Rem. Supp. 1945 § 6163-99.]

69.04.330 Food—Misbranding by failure to show artificial flavoring, coloring, etc. If a food bears or contains any artificial flavoring, artificial coloring, or chemical preservative, it shall be deemed to be misbranded unless it bears labeling stating that fact: PROVIDED, That to the extent that compliance with the requirements of this section is impracticable, exemptions shall be established by regulations promulgated by the director. The provisions of this section and of RCW 69.04.280 and 69.04.310, with respect to artificial coloring, shall not apply in the case of butter, cheese, or ice cream. [1945 c 257 § 51; Rem. Supp. 1945 § 6163-100.]

69.04.331 Popcorn sold by theaters or commercial food service establishments—Misbranded if the use of butter or ingredients of butter-like flavoring not disclosed. (1) If a theater or other commercial food service establishment prepares and sells popcorn for human consumption, the establishment, at the point of sale, shall disclose by posting a sign in a conspicuous manner to prospective consumers a statement as to whether the butter or butter-like flavoring added to or attributed to the popcorn offered for sale is butter or is some other product. If the flavoring is some other product, the establishment shall also disclose the ingredients of the product.

The director of agriculture shall adopt rules prescribing the size and content of the sign upon which the disclosure is to be made. Any popcorn sold by or offered for sale by such an establishment to a consumer in violation of this section or the provisions of RCW 69.04.280, it shall be deemed to be misbranded unless its label bears (1) the common or usual name of the food, if any there be, and (2) in case it is fabricated from two or more ingredients, the common or usual name of each such ingredient; except that spices, flavorings, and colorings, other than those sold as such, may be designated as spices, flavorings, and colorings without naming each: PROVIDED, That, to the extent that compliance with the requirements of clause (2) of this section is impracticable, or results in deception or unfair competition, exemptions shall be established by regulations promulgated by the director. [1945 c 257 § 49; Rem. Supp. 1945 § 6163-98.]
69.04.350 Permits to manufacture or process certain foods. Whenever the director finds after investigation that the distribution in intrastate commerce of any class of food may, by reason of contamination with micro-organisms during the manufacture, processing, or packing thereof in any locality, be injurious to health, and that such injurious nature cannot be adequately determined after such articles have entered intrastate commerce, he or she then, and in such case only, shall promulgate regulations providing for the issuance, to manufacturers, processors, or packers of such class of food in such locality, of permits to which shall be attached such conditions governing the manufacture, processing, or packing of such class of food, for such temporary period of time, as may be necessary to protect the public health; and after the effective date of such regulations, and during such temporary period, no person shall introduce or deliver for introduction into intrastate commerce, any such food manufactured, processed, or packed by any such manufacturer, processor, or packer unless such manufacturer, processor, or packer holds a permit issued by the director as provided by such regulations. Insofar as practicable, such regulations shall conform with, shall specify the conditions prescribed by, and shall remain in effect only so long as those promulgated under section 404(a) of the federal act. [2012 c 257 § 53; Rem. Supp. 1945 § 6163-103.]

69.04.333 Poultry and poultry products—Label to indicate if product frozen. It shall be unlawful for any person to sell at retail or display for sale at retail any poultry and poultry products, including turkey, which has been frozen at any time, without having the package or container in which the same is sold bear a label clearly discernible to a customer that such product has been frozen and whether or not the same has since been thawed. No such poultry or poultry product shall be sold unless in such a package or container bearing said label. [1969 ex.s. c 194 § 1.]

69.04.334 Turkeys—Label requirement as to grading. No person shall advertise for sale, sell, offer for sale or hold for sale in intrastate commerce any turkey that does not bear a label. Such label shall be properly displayed on the package if such turkey is prepackaged, or attached to the turkey if not prepackaged. Such label shall, if the turkey has been graded, state the name of the governmental agency, whether federal or state, and the grade. No turkey which has been graded may be labeled as being ungraded. Any advertisement in any media concerning the sale of turkeys shall state or set forth whether a turkey is ungraded or graded and the specific grade if graded. [1969 ex.s. c 194 § 2.]

69.04.335 RCW 69.04.333 and 69.04.334 subject to enforcement and penalty provisions of chapter. The provisions of this chapter shall be applicable to the enforcement of RCW 69.04.333 and 69.04.334 and any person violating the provisions of RCW 69.04.333 and 69.04.334 shall be subject to the applicable civil and criminal penalties for such violations as provided for in this chapter. [1969 ex.s. c 194 § 3.]

69.04.340 Natural vitamin, mineral, or dietary properties need not be shown. Nothing in this chapter shall be construed to require the labeling or advertising to indicate the natural vitamin, natural mineral, or other natural dietary properties of dairy products or other agricultural products when sold as food. [1945 c 257 § 52; Rem. Supp. 1945 § 6163-101.]

69.04.350 Permits to manufacture or process certain foods. Whenever the director finds after investigation that the distribution in intrastate commerce of any class of food may, by reason of contamination with micro-organisms during the manufacture, processing, or packing thereof in any locality, be injurious to health, and that such injurious nature cannot be adequately determined after such articles have
tion of RCW 69.04.210(2)(a); but when such substance is so required or cannot be so avoided, the director shall promulgate regulations limiting the quantity therein or thereon to such extent as he or she finds necessary for the protection of public health, and any quantity exceeding the limits so fixed shall also be deemed unsafe for purposes of the application of RCW 69.04.210(2)(a). While such a regulation is in effect limiting the quantity of any such substance in the case of any food, such food shall not, by reason of bearing or containing any added amount of such substance, be considered to be adulterated within the meaning of RCW 69.04.210(1). In determining the quantity of such added substance to be tolerated in or on different articles of food, the director shall take into account the extent to which the use of such substance is required or cannot be avoided in the production of each such article, and the other ways in which the consumer may be affected by the same or other poisonous or deleterious substances. [2012 c 117 § 336; 1963 c 198 § 2; 1945 c 257 § 57; Rem. Supp. 1945 § 6163-106.]

69.04.392 Regulations permitting tolerance of harmful matter—Pesticide chemicals in or on raw agricultural commodities. (1) Any poisonous or deleterious pesticide chemical, or any pesticide chemical which generally is recognized among experts qualified by scientific training and experience to evaluate the safety of pesticide chemicals as unsafe for use, added to a raw agricultural commodity, shall be deemed unsafe for the purpose of the application of RCW 69.04.210(2)(a) unless:

(a) A tolerance for such pesticide chemical in or on the raw agricultural commodity has been prescribed pursuant to subsection (2) of this section and the quantity of such pesticide chemical in or on the raw agricultural commodity is within the limits of the tolerance so prescribed; or

(b) With respect to use in or on such raw agricultural commodity, the pesticide chemical has been exempted from the requirement of a tolerance pursuant to subsection (2) of this section.

While a tolerance or exemption from tolerance is in effect for a pesticide chemical with respect to any raw agricultural commodity, such raw agricultural commodity shall not, by reason of bearing or containing any added amount of such pesticide chemical, be considered to be adulterated within the meaning of RCW 69.04.210(1).

(2) The regulations promulgated under section 408 of the federal food, drug and cosmetic act, as of July 1, 1975, setting forth the tolerances for pesticide chemicals in or on any raw agricultural commodity, are hereby adopted as the regulations for tolerances applicable to this chapter: PROVIDED, That the director is hereby authorized to adopt by regulation any new or future amendments to such federal regulations for tolerances, including exemption from tolerance and zero tolerances, to the extent necessary to protect the public health. The director is also authorized to issue regulations in the absence of federal regulations and to prescribe therein tolerances for pesticides, exemptions, and zero tolerances, upon his or her own motion or upon the petition of any interested party requesting that such a regulation be established. It shall be incumbent upon such petitioner to establish, by data submitted to the director, that a necessity exists for such regulation and that the effect of such regulation will not be detrimental to the public health. If the data furnished by the petitioner is not sufficient to allow the director to determine whether such a regulation should be promulgated, the director may require additional data to be submitted and failure to comply with this request shall be sufficient grounds to deny the request of the petitioner for the issuance of such regulation.

(3) In adopting any new or amended tolerances by regulation issued pursuant to this section, the director shall give appropriate consideration, among other relevant factors, to the following: (a) The purpose of this chapter being to promote uniformity of state legislation with the federal act; (b) the necessity for the production of an adequate, wholesome, and economical food supply; (c) the other ways in which the consumer may be affected by the same pesticide chemical or by other related substances that are poisonous or deleterious; and (d) the opinion of experts qualified by scientific training and experience to determine the proper tolerance to be allowed for any pesticide chemical. [2012 c 117 § 337; 1975 1st ex.s. c 7 § 26; 1963 c 198 § 3.]

Purpose of section: See RCW 69.04.398.

69.04.394 Regulations permitting tolerance of harmful matter—Food additives. (1) A food additive shall, with respect to any particular use or intended use of such additive, be deemed unsafe for the purpose of the application of clause (2)(c) of RCW 69.04.210, unless:

(a) It and its use or intended use conform to the terms of an exemption granted, pursuant to a regulation under subsection (2) hereof providing for the exemption from the requirements of this section for any food additive, and any food bearing or containing such additive, intended solely for investigational use by qualified experts when in the director’s opinion such exemption is consistent with the public health; or

(b) There is in effect, and it and its use or intended use are in conformity with a regulation issued or effective under subsection (2) hereof prescribing the conditions under which such additive may be safely used.

While such a regulation relating to a food additive is in effect, a food shall not, by reason of bearing or containing such an additive in accordance with the regulation, be considered adulterated within the meaning of clause (1) of RCW 69.04.210.

(2) The regulations promulgated under section 409 of the Federal Food, Drug and Cosmetic Act, as of July 1, 1975, prescribing the conditions under which such food additive may be safely used, are hereby adopted as the regulations applicable to this chapter: PROVIDED, That the director is hereby authorized to adopt by regulation any new or future amendments to the federal regulations. The director is also authorized to issue regulations in the absence of federal regulations and to prescribe the conditions under which a food additive may be safely used and exemptions where such food additive is to be used solely for investigational purposes; either upon his or her own motion or upon the petition of any interested party requesting that such a regulation be established. It shall be incumbent upon such petitioner to establish, by data submitted to the director, that a necessity exists for such regulation and that the effect of such a regulation will not be detrimental to the public health. If the data fur-
lished by the petitioner is not sufficient to allow the director to determine whether such a regulation should be promulgated, the director may require additional data to be submitted and failure to comply with this request shall be sufficient grounds to deny the request of the petitioner for the issuance of such a regulation.

(3) In adopting any new or amended regulations pursuant to this section, the director shall give appropriate consideration, among other relevant factors, to the following: (a) The purpose of this chapter being to promote uniformity of state legislation with the federal act; (b) the probable consumption of the additive and of any substance formed in or on food because of the use of the additive; (c) the cumulative effect of such additive in the diet of human beings or animals, taking into account any chemically or pharmacologically related substance or substances in such diet; and (d) safety factors which in the opinion of experts qualified by scientific training and experience to evaluate the safety of food additives are generally recognized as appropriate for the use of animal experimentation data. [2009 c 549 § 1021; 1975 1st ex.s. c 7 § 27; 1963 c 198 § 4.]

Purpose of section: See RCW 69.04.398.

69.04.396 Regulations permitting tolerance of harmful matter—Color additives. (1) A color additive shall, with respect to any particular use (for which it is being used or intended to be used or is represented as suitable) in or on food, be deemed unsafe for the purpose of the application of RCW 69.04.231, unless:

(a) There is in effect, and such color additive and such use are in conformity with, a regulation issued under this section listing such additive for such use, including any provision of such regulation prescribing the conditions under which such additive may be safely used;

(b) Such additive and such use thereof conform to the terms of an exemption for experimental use which is in effect pursuant to regulation under this section.

While there are in effect regulations under this section relating to a color additive or an exemption with respect to such additive a food shall not, by reason of bearing or containing such additive in all respects in accordance with such regulations or such exemption, be considered adulterated within the meaning of clause (1) of RCW 69.04.210.

(2) The regulations promulgated under section 706 of the Federal Food, Drug and Cosmetic Act, as of July 1, 1975, prescribing the use or limited use of such color additive, are hereby adopted as the regulations applicable to this chapter: PROVIDED, That the director is hereby authorized to adopt by regulation any new or future amendments to the federal regulations. The director is also authorized to issue regulations in the absence of federal regulations and to prescribe therein the conditions under which a color additive may be safely used including exemptions for experimental purposes. Such a regulation may be issued either upon the director’s own motion or upon the petition of any interested party requesting that such a regulation be established. It shall be incumbent upon such petitioner to establish, by data submitted to the director, that a necessity exists for such regulation and that the effect of such a regulation will not be detrimental to the public health. If the data furnished by the petitioner is not sufficient to allow the director to determine whether such a regulation should be promulgated, the director may require additional data to be submitted and failure to comply with this request shall be sufficient grounds to deny the request of the petitioner for the issuance of such a regulation.

(3) In adopting any new or amended regulations pursuant to this section, the director shall give appropriate consideration, among other relevant factors, to the following: (a) The purpose of this chapter being to promote uniformity of state legislation with the federal act; (b) the probable consumption of, or other relevant exposure from, the additive and of any substance formed in or on food because of the use of the additive; (c) the cumulative effect, if any, of such additive in the diet of human beings or animals, taking into account the same or any chemically or pharmacologically related substance or substances in such diet; (d) safety factors which, in the opinion of experts qualified by scientific training and experience to evaluate the safety of color additives for the use or uses for which the additive is proposed to be listed, are generally recognized as appropriate for the use of animal experimentation data; (e) the availability of any needed practicable methods of analysis for determining the identity and quantity of (i) the pure dye and all intermediates and other impurities contained in such color additives, (ii) such additive in or on any article of food, and (iii) any substance formed in or on such article because of the use of such additive; and (f) the conformity by the manufacturer with the established standards in the industry relating to the proper formation of such color additive so as to result in a finished product safe for use as a color additive. [2009 c 549 § 1022; 1975 1st ex.s. c 7 § 28; 1963 c 198 § 6.]

Purpose of section: See RCW 69.04.398.

Food—Adulteration by color additive: RCW 69.04.231.

69.04.410 Purpose of RCW 69.04.110, 69.04.392, 69.04.394, 69.04.396—Uniformity with federal laws and regulations—Application to production of kosher food products—Adoption of rules. (1) The purpose of RCW 69.04.110, 69.04.392, 69.04.394, and 69.04.396 is to promote uniformity of state legislation and rules with the Federal Food, Drug and Cosmetic Act 21 USC 301 et seq. and regulations adopted thereunder. In accord with such declared purpose any regulation adopted under said federal food, drug and cosmetic act concerning food in effect on July 1, 1975, and not adopted under any other specific provision of RCW 69.04.110, 69.04.392, 69.04.394, and 69.04.396 are hereby deemed to have been adopted under the provision hereof. Further, to promote such uniformity any regulation adopted hereafter under the provisions of the federal food, drug and cosmetic act concerning food and published in the federal register shall be deemed to have been adopted under the provisions of RCW 69.04.110, 69.04.392, 69.04.394, and 69.04.396 in accord with chapter 34.05 RCW as enacted or hereafter amended. The director may, however, within thirty days of the publication of the adoption of any such regulation under the federal food, drug and cosmetic act give public notice that a hearing will be held to determine if such regulation shall not be applicable under the provisions of RCW 69.04.110, 69.04.392, 69.04.394, and 69.04.396. Such hearing shall be in accord with the requirements of chapter 34.05 RCW as enacted or hereafter amended.
(2) The provisions of subsection (1) of this section do not apply to rules adopted by the director as necessary to permit the production of kosher food products as defined in RCW 69.90.010.

(3) Notwithstanding the provisions of subsections (1) and (2) of this section the director may adopt rules necessary to carry out the provisions of this chapter. [1991 c 162 § 5; 1986 c 203 § 18; 1975 1st ex.s. c 7 § 36.]

Additional notes found at www.leg.wa.gov

69.04.400 Conformance with federal regulations. The regulations promulgated under RCW 69.04.390 shall conform, insofar as practicable, with those promulgated under section 406 of the federal act. [1963 c 198 § 7; 1945 c 257 § 58; Rem. Supp. 1945 § 6163-107.]

69.04.410 Drugs—Adulteration by harmful substances. A drug or device shall be deemed to be adulterated (1) if it consists in whole or in part of any filthy, putrid, or decomposed substance; or (2) if it has been produced, prepared, packed, or held under insanitary conditions whereby it may have been contaminated with filth, or whereby it may have been rendered injurious to health; or (3) if it is a drug and its container is composed in whole or in part of any poisonous or deleterious substance which may render the contents injurious to health; or (4) if it is a drug and it bears or contains, for purposes of coloring only, a coal tar color other than one that is harmless and suitable for use in drugs for such purposes, as provided by regulations promulgated under section 504 of the federal act. [1945 c 257 § 61; Rem. Supp. 1945 § 6163-108. Prior: 1923 c 36 § 1; 1907 c 211 § 3; 1901 c 94 § 3.]

69.04.420 Drugs—Adulteration for failure to comply with compendium standard. If a drug or device purports to be or is represented as a drug the name of which is recognized in an official compendium, and its strength differs from, or its quality or purity falls below, the standard set forth in such compendium, it shall be deemed to be adulterated. Such determination as to strength, quality or purity shall be made in accordance with the tests or methods of assay set forth in such compendium or prescribed by regulations promulgated under section 501(b) of the federal act. No drug defined in an official compendium shall be deemed to be adulterated under this section because it differs from the standard of strength, quality, or purity therefor set forth in such compendium, if its difference in strength, quality, or purity from such standard is plainly stated on its label. Whenever a drug is recognized in both the United States pharmacopoeia and the homeopathic pharmacopoeia of the United States, it shall be subject to the requirements of the United States pharmacopoeia unless it is labeled and offered for sale as a homeopathic drug, in which case it shall be subject to the provisions of the homeopathic pharmacopoeia of the United States and not to those of the United States pharmacopoeia. [1945 c 257 § 60; Rem. Supp. 1945 § 6163-109.]

69.04.430 Drugs—Adulteration for lack of represented purity or quality. If a drug or device is not subject to the provisions of RCW 69.04.420 and its strength differs from, or its purity or quality falls below, that which it purports or is represented to possess, it shall be deemed to be adulterated. [1945 c 257 § 61; Rem. Supp. 1945 § 6163-110.]

69.04.440 Drugs—Adulteration by admixture or substitution of ingredients. A drug shall be deemed to be adulterated if any substance has been (1) mixed or packed therewith so as to reduce its quality or strength or (2) substituted wholly or in part therefor. [1945 c 257 § 62; Rem. Supp. 1945 § 6163-111.]

69.04.450 Drugs—Misbranding by false labeling. A drug or device shall be deemed to be misbranded if its labeling is false or misleading in any particular. [1945 c 257 § 63; Rem. Supp. 1945 § 6163-112. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.460 Packaged drugs—Misbranding. If a drug or device is in package form, it shall be deemed to be misbranded unless it bears a label containing (1) the name and place of business of the manufacturer, packer, or distributor; and (2) an accurate statement of the quantity of the contents in terms of weight, measure, or numerical count: PROVIDED, That under clause (2) of this section reasonable variations shall be permitted, and exemptions as to small packages shall be established, by regulations promulgated by the director. [1945 c 257 § 64; Rem. Supp. 1945 § 6163-113. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.470 Drugs—Misbranding by lack of prominent label. A drug or device shall be deemed to be misbranded if any word, statement, or other information required by or under authority of this chapter to appear on the label or labeling is not prominently placed thereon with such conspicuousness (as compared with other words, statements, designs, or devices, in the labeling) and in such terms as to render it likely to be read and understood by the ordinary individual under customary conditions of purchase and use. [1945 c 257 § 65; Rem. Supp. 1945 § 6163-114. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.480 Drugs—Misbranding for failure to state content of habit forming drug. A drug or device shall be deemed to be misbranded if it is for use by human beings and contains any quantity of the narcotic or hypnotic substance alpha eucaine, barbituric acid, beta eucaine, bromal, cannab is, carbromal, chloral, cocoa, cocaine, codeine, heroin, marijuana, morphine, opium, paraldehyde, peyote, or sulpho methane; or any chemical derivative of such substance, which derivative has been designated as habit forming by regulations promulgated under section 502(d) of the federal act; unless its label bears the name and quantity or proportion of such substance or derivative and in juxtaposition therewith the statement "Warning—May be habit forming." [2009 c 549 § 1023; 1945 c 257 § 66; Rem. Supp. 1945 § 6163-115. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.490 Drugs—Misbranding by failure to show usual name and ingredients. If a drug is not designated solely by a name recognized in an official compendium it shall be deemed to be misbranded unless its label bears (1) the common or usual name of the drug, if such there be; and
(2), in case it is fabricated from two or more ingredients, the common or usual name of each active ingredient, including the quantity, kind, and proportion of any alcohol, and also including, whether active or not, the name and quantity or proportion of any bromides, ether, chloroform, acetanilid, acethypetidin, amidopyrine, antipyrine, atropine, hyoscine, hyoscynamine, arsenic, digitalis, glucosides, mercury, ouabain, strophanthidin, strychnine, thyroid, or any derivative or preparation of any such substances, contained therein: PROVIDED, That to the extent that compliance with the requirements of clause (2) of this section is impracticable, exemptions shall be established by regulations promulgated by the director. [1945 c 257 § 67; Rem. Supp. 1945 § 6163-116. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.500 Drugs—Misbranding by failure to give directions for use and warnings. A drug or device shall be deemed to be misbranded unless its labeling bears (1) adequate directions for use; and (2) such adequate warnings against use in those pathological conditions or by children where its use may be dangerous to health, or against unsafe dosage or methods or duration of administration or application, in such manner and form, as are necessary for the protection of users: PROVIDED, That where any requirement of clause (1) of this section as applied to any drug or device, is not necessary for the protection of the public health, the director shall promulgate regulations exempting such drug or device from such requirements. Such regulations shall include the exemptions prescribed under section 502(f)(1) of the federal act, insofar as such exemptions are applicable hereunder. [1945 c 257 § 68; Rem. Supp. 1945 § 6163-117. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.510 Drugs—Misbranding for improper packaging and labeling. A drug or device shall be deemed to be misbranded if it purports to be a drug the name of which is recognized in an official compendium, unless it is packaged and labeled as prescribed therein: PROVIDED, That the method of packing may be modified with the consent of the director, as permitted under section 502(g) of the federal act. Whenever a drug is recognized in both the United States pharmacopoeia and the homeopathic pharmacopoeia of the United States, it shall be subject to the requirements of the United States pharmacopoeia with respect to packaging and labeling unless it is labeled and offered for sale as a homeopathic drug, in which case it shall be subject to the provisions of the homeopathic pharmacopoeia of the United States, and not to those of the United States pharmacopoeia. [1945 c 257 § 69; Rem. Supp. 1945 § 6163-118. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.520 Drugs—Misbranding for failure to show possibility of deterioration. If a drug or device has been found by the secretary of agriculture of the United States to be a drug liable to deterioration, it shall be deemed to be misbranded unless it is packaged in such form and manner, and its label bears a statement of such precautions, as required in an official compendium or by regulations promulgated under section 502(h) of the federal act for the protection of the public health. [1945 c 257 § 70; Rem. Supp. 1945 § 6163-119. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.530 Drugs—Misbranding by misleading representation. A drug shall be deemed to be misbranded if (1) its container is so made, formed, or filled as to be misleading; or (2) if it is an imitation of another drug; or (3) if it is offered for sale under the name of another drug; or (4) if it is dangerous to health when used in the dosage, or with the frequency or duration prescribed, recommended, or suggested in the labeling thereof. [1945 c 257 § 71; Rem. Supp. 1945 § 6163-120. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.540 Drugs—Misbranding by sale without prescription of drug requiring it. A drug or device shall be deemed to be misbranded if it is a drug which by label provides, or which the federal act or any applicable law requires by label to provide, in effect, that it shall be used only upon the prescription of a physician, dentist, or veterinarian, unless it is dispensed at retail on a written prescription signed by a physician, dentist, or veterinarian, who is licensed by law to administer such a drug. [1945 c 257 § 72; Rem. Supp. 1945 § 6163-121. Prior: 1923 c 36 § 2; 1907 c 211 § 4.]

69.04.550 Drugs exempt if in transit for completion purposes. A drug or device which is, in accordance with the practice of the trade, to be processed, labeled, or repacked in substantial quantities at an establishment other than the establishment where it was originally processed or packed, is exempted from the affirmative labeling and packaging requirements of this chapter, while it is in transit in intrastate commerce from the one establishment to the other, if such transit is made in good faith for such completion purposes only; but it is otherwise subject to all the applicable provisions of this chapter. [1945 c 257 § 73; Rem. Supp. 1945 § 6163-122.]

69.04.560 Dispensing of certain drugs exempt. A drug dispensed on a written prescription signed by a physician, dentist, or veterinarian (except a drug dispensed in the course of the conduct of a business of dispensing drugs pursuant to diagnosis by mail) shall, if (1) such physician, dentist, or veterinarian is licensed by law to administer such drug, and (2) such drug bears a label containing the name and place of business of the dispenser, the serial number and date of such prescription, and the name of such physician, dentist, or veterinarian, be exempt from the requirements of RCW 69.04.450 through 69.04.540. [1945 c 257 § 74; Rem. Supp. 1945 § 6163-123.]

69.04.565 DMSO (dimethyl sulfoxide) authorized. Notwithstanding any other provision of state law, DMSO (dimethyl sulfoxide) may be introduced into intrastate commerce as long as (1) it is manufactured or distributed by persons licensed pursuant to chapter 18.64 RCW or chapter 18.92 RCW, and (2) it is used, or intended to be used, in the treatment of human beings or animals for any ailment or adverse condition: PROVIDED, That DMSO intended for topical application, consistent with rules governing purity and labeling promulgated by the state board of pharmacy, shall not be considered a legend drug and may be sold by any retailer. [1981 c 50 § 1.]
69.04.570 Introduction of new drug. No person shall introduce or deliver for introduction into intrastate commerce any new drug which is subject to section 505 of the federal act unless an application with respect to such drug has become effective thereunder. No person shall introduce or deliver for introduction into intrastate commerce any new drug which is not subject to section 505 of the federal act, unless (1) it has been found, by appropriate tests, that such drug is not unsafe for use under the conditions prescribed, recommended, or suggested in the labeling thereof; and (2) an application has been filed under this section of this chapter with respect to such drug: PROVIDED, That the requirement of subsection (2) of this section shall not apply to any drug introduced into intrastate commerce at any time prior to the enactment of this chapter or into interstate commerce at any time prior to the enactment of the federal act: PROVIDED FURTHER, That if the director finds that the requirement of subsection (2) of this section as applied to any drug or class of drugs, is not necessary for the protection of the public health, he or she shall promulgate regulations of exemption accordingly. [1945 c 257 § 79; Rem. Supp. 1945 § 6163-124.]

69.04.580 Application for introduction. An application under RCW 69.04.570 shall be filed with the director, and subject to any waiver by the director, shall include (1) full reports of investigations which have been made to show whether or not the drug, subject to the application, is safe for use under the conditions prescribed, recommended, or suggested in the labeling thereof; (2) a full list of the articles used as components of such drug; (3) a full statement of the composition of such drug; (4) a full description of the methods used in, and the facilities and controls used for, the manufacture, processing, and packing of such drug; (5) such samples of such drug and of the articles used as components thereof as the director may require; and (6) specimens of the labeling proposed to be used for such drug. [1945 c 257 § 76; Rem. Supp. 1945 § 6163-125.]

69.04.590 Effective date of application. An application filed under RCW 69.04.570 shall become effective on the sixtieth day after the filing thereof, unless the director (1) makes such application effective prior to such day; or (2) issues an order with respect to such application pursuant to RCW 69.04.600. [1945 c 257 § 77; Rem. Supp. 1945 § 6163-126.]

69.04.600 Denial of application. If the director finds, upon the basis of the information before him or her and after due notice and opportunity for hearing to the applicant, that the drug, subject to the application, is not safe for use under the conditions prescribed, recommended, or suggested in the labeling thereof, he or she shall, prior to such effective date, issue an order refusing to permit such application to become effective and stating the findings upon which it is based. [1945 c 257 § 78; Rem. Supp. 1945 § 6163-127.]

69.04.610 Revocation of denial. An order refusing to permit an application under RCW 69.04.570 to become effective may be suspended or revoked by the director, for cause and by order stating the findings upon which it is based. [1945 c 257 § 79; Rem. Supp. 1945 § 6163-128.]

69.04.620 Service of order of denial. Orders of the director issued under RCW 69.04.600 shall be served (1) in person by a duly authorized representative of the director or (2) by mailing the order by registered mail addressed to the applicant or respondent at his or her address last known to the director. [2012 c 117 § 340; 1945 c 257 § 80; Rem. Supp. 1945 § 6163-129.]

69.04.630 Drug for investigational use exempt. A drug shall be exempt from the operation of RCW 69.04.570 which is intended, and introduced or delivered for introduction into intrastate commerce, solely for investigational use by experts qualified by scientific training and experience to investigate the safety of drugs and which is plainly labeled "For investigational use only." [1945 c 257 § 81; Rem. Supp. 1945 § 6163-130.]

69.04.640 Court review of denial. The superior court of Thurston county shall have jurisdiction to review and to affirm, modify, or set aside any order issued under RCW 69.04.600, upon petition seasonably made by the person to whom the order is addressed and after prompt hearing upon due notice to both parties. [1945 c 257 § 82; Rem. Supp. 1945 § 6163-131.]

69.04.650 Dispensing of certain drugs exempt. A drug dispensed on a written prescription signed by a physician, dentist, or veterinarian (except a drug dispensed in the course of the conduct of a business of dispensing drugs pursuant to diagnosis by mail) shall, if (1) such physician, dentist, or veterinarian is licensed by law to administer such drug, and (2) such drug bears a label containing the name and place of business of the dispenser, the serial number and date of such prescription, and the name of such physician, dentist, or veterinarian, be exempt from the operation of RCW 69.04.570 through 69.04.640. [1945 c 257 § 83; Rem. Supp. 1945 § 6163-132.]

69.04.660 Federally licensed drugs exempt. The provisions of RCW 69.04.570 shall not apply to any drug which is licensed under the federal virus, serum, and toxin act of July 1, 1902; or under the federal virus, serums, toxins, anti-toxins, and analogous products act of March 4, 1913. [1945 c 257 § 84; Rem. Supp. 1945 § 6163-133.]

69.04.670 Cosmetics—Adulteration by injurious substances. A cosmetic shall be deemed to be adulterated (1) if it bears or contains any poisonous or deleterious substance which may render it injurious to users under the conditions of use prescribed in the labeling thereof, or under such conditions of use as are customary or usual: PROVIDED, That this provision shall not apply to coal tar hair dye, the label of which bears the following legend conspicuously displayed thereon: "Caution—This product contains ingredients which may cause skin irritation on certain individuals and a preliminary test according to accompanying directions should first be made. This product must not be used for dyeing the eyelashes or eyebrows; to do so may cause blindness.", and
the labeling of which bears adequate directions for such preliminary testing. For the purposes of this paragraph and paragraph (5) the term "hair dye" shall not include eyelash dyes or eyebrow dyes; or (2) if it consists in whole or in part of any filthy, putrid, or decomposed substance; or (3) if it has been produced, prepared, packed, or held under insanitary conditions whereby it may have become contaminated with filth, or whereby it may have been rendered injurious to health; or (4) if its container is composed in whole or in part of any poisonous or deleterious substance which may render the contents injurious to health; or (5) if it is not a hair dye and it bears or contains a coal tar color other than one that is harmless and suitable for use in cosmetics, as provided by regulations promulgated under section 604 of the federal act. [1945 c 257 § 85; Rem. Supp. 1945 § 6163-134.]

69.04.680 Cosmetics—Misbranding by false label, etc. A cosmetic shall be deemed to be misbranded (1) if its labeling is false or misleading in any particular; or (2) if in package form, unless it bears a label containing (a) the name and place of business of the manufacturer, packer, or distributor; and (b) an accurate statement of the quantity of the contents in terms of weight, measure, or numerical count: PROVIDED, That under clause (b) of this section reasonable variations shall be permitted, and exemptions as to small packages shall be established, by regulations prescribed by the director. [1945 c 257 § 86; Rem. Supp. 1945 § 6163-135.]

69.04.690 Cosmetics—Misbranding by lack of prominent label. A cosmetic shall be deemed to be misbranded (1) if any word, statement, or other information required by or under authority of this chapter to appear on the label or labeling is not prominently placed thereon with such conspicuously (as compared with other words, statements, designs, or devices, in the labeling) and in such terms as to render it likely to be read and understood by the ordinary individual under customary conditions of purchase and use; or (2) if its container is so made, formed, or filled as to be misleading. [1945 c 257 § 87; Rem. Supp. 1945 § 6163-136.]

69.04.700 Cosmetics exempt if in transit for completion purposes. A cosmetic which is, in accordance with the practice of the trade, to be processed, labeled, or repacked in substantial quantities at an establishment other than the establishment where it was originally processed or packed, is exempted from the affirmative labeling requirements of this chapter, while it is in transit in intrastate commerce from the one establishment to the other, if such transit is made in good faith for such completion purposes only; but it is otherwise subject to all the applicable provisions of this chapter. [1945 c 257 § 88; Rem. Supp. 1945 § 6163-137.]

69.04.710 Advertisement, when deemed false. An advertisement of a food, drug, device, or cosmetic shall be deemed to be false, if it is false or misleading in any particular. [1945 c 257 § 89; Rem. Supp. 1945 § 6163-138.]

69.04.720 Advertising of cure of certain diseases deemed false. The advertisement of a drug or device representing it to have any effect in albuminuria, appendicitis, arteriosclerosis, blood poison, bone disease, Bright's disease, cancer, carbuncles, cholecystitis, diabetes, diphtheria, dropsy, erysipelas, gallstones, heart and vascular diseases, high blood pressure, mastoiditis, measles, meningitis, mumps, nephritis, otitis media, paralysis, pneumonia, poliomyelitis (infantile paralysis), prostate gland disorders, pyelitis, scarlet fever, sexual impotence, sinus infection, smallpox, tuberculosis, tumors, typhoid, uremia, *venereal disease, shall also be deemed to be false; except that no advertisement not in violation of RCW 69.04.710 shall be deemed to be false under this section if it is disseminated only to members of the medical, veterinary, dental, pharmacal, and other legally recognized professions dealing with the healing arts, or appears only in the scientific periodicals of these professions, or is disseminated only for the purpose of public health education by persons not commercially interested, directly or indirectly, in the sale of such drugs or devices: PROVIDED, That whenever the director determines that an advance in medical science has made any type of self-medication safe as to any of the diseases named above, the director shall by regulation authorize the advertisement of drugs having curative or therapeutic effect for such disease, subject to such conditions and restrictions as the director may deem necessary in the interest of public health: PROVIDED FURTHER, That this section shall not be construed as indicating that self-medication for diseases other than those named herein is safe or efficacious. [1945 c 257 § 90; Rem. Supp. 1945 § 6163-139.]

*Reviser's note: The term "venereal disease" was changed to "sexually transmitted disease" by 1988 c 206.

69.04.730 Enforcement, where vested—Regulations. The authority to promulgate regulations for the efficient enforcement of this chapter is hereby vested in the director: PROVIDED, HOWEVER, That the director shall designate the Washington state board of pharmacy to carry out all the provisions of this chapter pertaining to drugs and cosmetics, with authority to promulgate regulations for the efficient enforcement thereof. [1945 c 257 § 91 (vetoed); 1947 c 25 (passed notwithstanding veto); Rem. Supp. 1947 § 6163-139a.]

69.04.740 Regulations to conform with federal regulations. The purpose of this chapter being to promote uniformity of state legislation with the federal act, the director is hereby authorized (1) to adopt, insofar as applicable, the regulations from time to time promulgated under the federal act; and (2) to make the regulations promulgated under this chapter conform, insofar as practicable, with those promulgated under the federal act. [1945 c 257 § 92; Rem. Supp. 1945 § 6163-140.]

69.04.750 Hearings. Hearings authorized or required by this chapter shall be conducted by the director or his or her duly authorized representative designated for the purpose. [2012 c 117 § 341; 1945 c 257 § 93; Rem. Supp. 1945 § 6163-141.]

69.04.761 Hearing on proposed regulation—Procedure. The director shall hold a public hearing upon a proposal to promulgate any new or amended regulation under this chapter. The procedure to be followed concerning such
hearings shall comply in all respects with chapter 34.05 RCW (Administrative Procedure Act) as now enacted or hereafter amended. [1963 c 198 § 13.]

**69.04.770 Review on petition prior to effective date.** The director shall have jurisdiction to review and to affirm, modify, or set aside any order issued under *RCW 69.04.760,* promulgating a new or amended regulation under this chapter, upon petition made at any time prior to the effective date of such regulation, by any person adversely affected by such order. [1945 c 257 § 95; Rem. Supp. 1945 § 6163-143.]

*Reviser’s note:* RCW 69.04.760 was repealed by 1963 c 198 § 15. Later enactment, see RCW 69.04.761.

**69.04.780 Investigations—Samples—Right of entry—Verified statements.** The director shall cause the investigation and examination of food, drugs, devices, and cosmetics subject to this chapter. The director shall have the right (1) to take a sample or specimen of any such article, for examination under this chapter, upon tendering the market price therefor to the person having such article in custody; and (2) to enter any place or establishment within this state, at reasonable times, for the purpose of taking a sample or specimen of any such article, for such examination.

The director and the director’s deputies, assistants, and inspectors are authorized to do all acts and things necessary to carry out the provisions of this chapter, including the taking of verified statements. Such department personnel are empowered to administer oaths of verification on the statements. [1991 c 162 § 6; 1945 c 257 § 96; Rem. Supp. 1945 § 6163-144.]

**69.04.790 Owner may obtain part of sample.** Where a sample or specimen of any such article is taken for examination under this chapter, the director shall, upon request, provide a part thereof for examination by any person named on the label of such article, or the owner thereof, or his or her attorney or agent; except that the director is authorized, by regulation, to make such reasonable exceptions from, and to impose such reasonable terms and conditions relating to, the operation of this section as he or she finds necessary for the proper administration of the provisions of this chapter. [2012 c 117 § 342; 1945 c 257 § 97; Rem. Supp. 1945 § 6163-145.]

**69.04.800 Access to records of other agencies.** For the purpose of enforcing the provisions of this chapter, pertinent records of any administrative agency of the state government shall be open to inspection by the director. [1945 c 257 § 98; Rem. Supp. 1945 § 6163-146.]

**69.04.810 Access to records of intrastate carriers.** For the purpose of enforcing the provisions of this chapter, carriers engaged in intrastate commerce, and persons receiving food, drugs, devices, or cosmetics in intrastate commerce or holding such articles so received, shall, upon the request of the director, permit the director at reasonable times, to have access to and to copy all records showing the movement in intrastate commerce of any food, drug, device, or cosmetic, or the holding thereof during or after such movement, and the quantity, shipper, and consignee thereof; and it shall be unlawful for any such carrier or person to fail to permit such access to and the copying of any such records so requested when such request is accompanied by a statement in writing specifying the nature or kind of food, drug, device, or cosmetic to which such request relates: PROVIDED, That evidence obtained under this section shall not be used in a criminal prosecution of the person from whom obtained: PROVIDED FURTHER, That except for violations of RCW 69.04.955, penalties levied under RCW 69.04.980, the requirements of RCW 69.04.950 through 69.04.980, and the requirements of this section, carriers shall not be subject to the other provisions of this chapter by reason of their receipt, carriage, holding, or delivery of food, drugs, devices, or cosmetics in the usual course of business as carriers. [1990 c 202 § 9; 1945 c 257 § 99; Rem. Supp. 1945 § 6163-147.]

**69.04.820 Right of entry to factories, warehouses, vehicles, etc.** For the purpose of enforcing the provisions of this chapter, the director is authorized (1) to enter, at reasonable times, any factory, warehouse, or establishment subject to this chapter, or to enter any vehicle being used to transport or hold food, drugs, devices, or cosmetics in intrastate commerce; and (2) to inspect, at reasonable times, such factory, warehouse, establishment, or vehicle and all pertinent equipment, finished and unfinished materials, containers, labeling, and advertisements therein. [1945 c 257 § 100; Rem. Supp. 1945 § 6163-148.]

**69.04.830 Publication of reports of judgments, orders and decrees.** The director may cause to be published from time to time reports summarizing all judgments, decrees, and court orders which have been rendered under this chapter, including the nature of the charge and the disposition thereof. [1945 c 257 § 101; Rem. Supp. 1945 § 6163-149.]

**69.04.840 Dissemination of information.** The director may cause to be disseminated information regarding food, drugs, devices, or cosmetics in situations involving, in the opinion of the director, imminent danger to health or gross deception of, or fraud upon, the consumer. Nothing in this section shall be construed to prohibit the director from collecting, reporting, and illustrating the results of his or her examinations and investigations under this chapter. [2012 c 117 § 343; 1945 c 257 § 102; Rem. Supp. 1945 § 6163-150.]

**69.04.845 Severability—1945 c 257.** If any provision of this chapter is declared unconstitutional, or the applicability thereof to any person or circumstances is held invalid, the constitutionality of the remainder of the chapter and the applicability thereof to other persons and circumstances shall not be affected thereby. [1945 c 257 § 103; Rem. Supp. 1945 § 6163-151.]

**69.04.850 Construction—1945 c 257.** This chapter and the regulations promulgated hereunder shall be so interpreted and construed as to effectuate its general purpose to secure uniformity with federal acts and regulations relating to adulterating, misbranding and false advertising of food, drugs, devices, and cosmetics. [1945 c 257 § 104; Rem. Supp. 1945 § 6163-152.]
69.04.860 Effective date of chapter—1945 c 257. This chapter shall take effect ninety days after the date of its enactment, and all state laws or parts of laws in conflict with this chapter are then repealed: PROVIDED, That the provisions of section 91 shall become effective on the enactment of this chapter, and thereafter the director is hereby authorized to conduct hearings and to promulgate regulations which shall become effective on or after the effective date of this chapter as the director shall direct: PROVIDED FURTHER, That all other provisions of this chapter to the extent that they may relate to the enforcement of such sections, shall take effect on the date of the enactment of this chapter. [1945 c 257 § 105; Rem. Supp. 1945 § 6163-153.]

Reviser’s note: 1945 c 257 § 91 referred to herein was vetoed by the governor but was subsequently reenacted as 1947 c 25 notwithstanding the veto. Section 91 is codified as RCW 69.04.730. For effective date of section 91 see preface 1947 session laws.

69.04.870 Short title. This chapter may be cited as the Uniform Washington Food, Drug, and Cosmetic Act. [1945 c 257 § 1; Rem. Supp. 1945 § 6163-50.]

69.04.880 Civil penalty. Whenever the director finds that a person has committed a violation of a provision of this chapter, the director may impose upon and collect from the violator a civil penalty not exceeding one thousand dollars per violation per day. Each and every such violation shall be a separate and distinct offense. Imposition of the civil penalty shall be subject to a hearing in conformance with chapter 34.05 RCW. [1991 c 162 § 2.]

69.04.900 Perishable packaged food—Pull date labeling—Definitions. For the purpose of RCW 69.04.900 through 69.04.920:

1) "Perishable packaged food goods" means and includes all foods and beverages, except alcoholic beverages, frozen foods, fresh meat, poultry and fish and a raw agricultural commodity as defined in this chapter, intended for human consumption which are canned, bottled, or packaged other than at the time and point of retail sale, which have a high risk of spoilage within a period of thirty days, and as determined by the director of the department of agriculture by rule and regulation to be perishable.

2) "Pull date" means the latest date a packaged food product shall be offered for sale to the public.

3) "Shelf life" means the length of time during which a packaged food product will retain its safe consumption quality if stored under proper temperature conditions.

4) "Fish" as used in subsection (1) of this section shall mean any water breathing animals, including, but not limited to, shellfish such as lobster, clams, crab, or other mollusca which are prepared, processed, sold, or intended or offered for sale. [1974 ex.s. c 57 § 1; 1973 1st ex.s. c 112 § 1.]

69.04.905 Perishable packaged food—Pull date labeling—Required. All perishable packaged food goods with a projected shelf life of thirty days or less, which are offered for sale to the public after January 1, 1974 shall state on the package the pull date. The pull date must be stated in day, and month and be in a style and format that is readily decipherable by consumers: PROVIDED, That the director of the department of agriculture may exclude the monthly requirement on the pull date for perishable packaged food goods which have a shelf life of seven days or less. No perishable packaged food goods shall be offered for sale after the pull date, except as provided in RCW 69.04.910. [1974 ex.s. c 57 § 2; 1973 1st ex.s. c 112 § 2.]

69.04.910 Perishable packaged food—Pull date labeling—Selling or trading goods beyond pull date—Repackaging to substitute for original date—Exception. No person shall sell, trade or barter any perishable packaged food goods beyond the pull date appearing thereon, nor shall any person rewrap or repack any perishable packaged food goods with the intention of placing a pull date thereon which is different from the original: PROVIDED, HOWEVER, That those packaged perishable food goods whose pull dates have expired may be sold if they are still wholesome and are without danger to health, and are clearly identified as having passed the pull date. [1973 1st ex.s. c 112 § 3.]

69.04.915 Perishable packaged food—Pull date labeling—Storage—Rules and regulations. The director of the department of agriculture shall by rule and regulation establish uniform standards for pull date labeling, and optimum storage conditions of perishable packaged food goods. In addition to his or her other duties, the director, in consultation with the secretary of the department of health where appropriate, may promulgate such other rules and regulations as may be necessary to carry out the purposes of RCW 69.04.900 through 69.04.920. [2012 c 117 § 344; 1989 1st ex.s. c 9 § 225; 1973 1st ex.s. c 112 § 4.]

Additional notes found at www.leg.wa.gov

69.04.920 Perishable packaged food—Pull date labeling—Penalties. Any person convicted of a violation of RCW 69.04.905 or 69.04.910 shall be punishable by a fine not to exceed five hundred dollars. [1973 1st ex.s. c 112 § 5.]

69.04.928 Seafood labeling requirements—Pamphlet—Direct retail endorsement. The department of agriculture must develop a pamphlet that generally describes the labeling requirements for seafood, as set forth in this chapter, and provide an adequate quantity of the pamphlets to the department of fish and wildlife to distribute with the issuance of a direct retail endorsement under RCW 77.65.510. [2002 c 301 § 11.]

Finding—Effective date—2002 c 301: See notes following RCW 77.65.510.

69.04.930 Frozen fish and meat—Labeling requirements—Exceptions. It shall be unlawful for any person to sell at retail or display for sale at retail any food fish as defined in RCW 77.08.022 or shellfish as defined in RCW 77.08.010, any meat, or any meat food product which has been frozen at any time, without having the package or container in which the same is sold bear a label clearly discernible to a customer that such product has been frozen and whether or not the same has since been thawed. No such food fish or shellfish, meat or meat food product shall be sold unless in such a package or container bearing said label: PROVIDED, That this section shall not include any of the aforementioned food or food products that have been frozen.
prior to being smoked, cured, cooked or subjected to the heat of commercial sterilization. [2003 c 39 § 28; 1999 c 291 § 32; 1988 c 254 § 8; 1983 1st ex.s. c 46 § 179; 1975 c 39 § 1.]

69.04.932 Salmon labeling—Definitions. Unless the context clearly requires otherwise, the definitions in this section apply throughout RCW 69.04.933 through 69.04.935.

(1) "Salmon" means all species of the genus Oncorhynchus, except those classified as game fish in Title 77 RCW, and includes:

<table>
<thead>
<tr>
<th>SCIENTIFIC NAME</th>
<th>COMMON NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oncorhynchus tsawtytscha</td>
<td>Chinook salmon or king salmon</td>
</tr>
<tr>
<td>Oncorhynchus kisutch</td>
<td>Coho salmon or silver salmon</td>
</tr>
<tr>
<td>Oncorhynchus keta</td>
<td>Chum salmon</td>
</tr>
<tr>
<td>Oncorhynchus gorbuscha</td>
<td>Pink salmon</td>
</tr>
<tr>
<td>Oncorhynchus nerka</td>
<td>Sockeye salmon</td>
</tr>
<tr>
<td>Salmo salar (in other than its landlocked form)</td>
<td>Atlantic salmon</td>
</tr>
</tbody>
</table>

(2) "Commercially caught" means salmon harvested by commercial fishers. [1993 c 282 § 2.]

Finding—1993 c 282: "The legislature finds that salmon consumers in Washington benefit from knowing the species and origin of the salmon they purchase. The accurate identification of such species, as well as knowledge of the country or state of origin and of whether they were caught commercially or were farm-raised, is important to consumers." [1993 c 282 § 1.]

69.04.933 Salmon labeling—Identification of species—Exceptions—Penalty. With the exception of a commercial fisher engaged in sales of fish to a fish buyer, no person may sell at wholesale or retail any fresh or frozen salmon food fish or cultured aquatic salmon without identifying the species of salmon by its common name to the buyer at the point of sale such that the buyer can make an informed decision in purchasing. A person knowingly violating this section is guilty of misbranding. This section shall not apply to salmon that is minced, pulverized, coated with batter, or breaded. [1993 c 282 § 3.]

Finding—1993 c 282: See note following RCW 69.04.932.

69.04.934 Salmon labeling—Identification as farm-raised or commercially caught—Exceptions—Penalty. With the exception of a commercial fisher engaged in sales of fish to a fish buyer, no person may sell at wholesale or retail any fresh or frozen:

(1) Private sector cultured aquatic salmon without identifying the product as farm-raised salmon; or

(2) Commercially caught salmon designated as food fish under Title 77 RCW without identifying the product as commercially caught salmon.

Identification of the products under subsections (1) and (2) of this section shall be made to the buyer at the point of sale such that the buyer can make an informed decision in purchasing.

A person knowingly violating this section is guilty of misbranding under this chapter. A person who receives misleading or erroneous information about whether the salmon is farm-raised or commercially caught, and subsequently inaccurately identifies salmon shall not be guilty of misbranding. This section shall not apply to salmon that is minced, pulverized, coated with batter, or breaded. [2003 c 39 § 29; 1993 c 282 § 4.]

Finding—1993 c 282: See note following RCW 69.04.932.

69.04.935 Salmon labeling—Rules for identification and enforcement. To promote honesty and fair dealing for consumers, the director, in consultation with the director of the department of fish and wildlife, shall adopt rules:

(1) Fixing and establishing a reasonable definition and standard of identity for salmon for purposes of identifying and selling salmon;

(2) Enforcing RCW 69.04.933 and 69.04.934. [1994 c 264 § 39; 1993 c 282 § 5.]

Finding—1993 c 282: See note following RCW 69.04.932.

69.04.940 Imported lamb products—Labeling requirements. All retail sales of fresh or frozen lamb products which are imported from another country shall be labelled with the country of origin. For the purposes of this section "imported lamb products" shall include but not be limited to, live lambs imported from another country but slaughtered in the United States. [1987 c 393 § 25.]

69.04.950 Transport of bulk foods—Definitions. The definitions in this section apply throughout RCW 69.04.950 through 69.04.980:

(1) "Food" means: (a) Any article used for food or drink for humans or used as a component of such an article; or (b) a food grade substance.

(2) "Food grade substance" means a substance which satisfies the requirements of the federal food, drug, and cosmetic act, meat inspection act, and poultry products act and rules promulgated thereunder as materials approved by the federal food and drug administration, United States department of agriculture, or United States environmental protection agency for use: (a) As an additive in food or drink for human consumption, (b) in sanitizing food or drink for human consumption, (c) in processing food or drink for human consumption, or (d) in maintaining equipment with food contact surfaces during which maintenance the substance is expected to come in contact with food or drink for human consumption.

(3) "In bulk form" means a food or substance which is not packaged or contained by anything other than the cargo carrying portion of the vehicle or vessel.

(4) "Vehicle or vessel" means a commercial vehicle or commercial vessel which has a gross weight of more than ten thousand pounds, is used to transport property, and is a motor vehicle, motor truck, trailer, railroad car, or vessel. [1990 c 202 § 1.]

Additional notes found at www.leg.wa.gov

69.04.955 Transport of bulk foods—Prohibitions—Exemption. (1) Except as provided in RCW 69.04.965 and 69.04.975, no person may transport in intrastate commerce food in bulk form in the cargo carrying portion of a vehicle or vessel that has been used for transporting in bulk form a cargo other than food.

(2) No person may transport in intrastate commerce food in bulk form in the cargo carrying portion of a vehicle or vessel unless the vehicle or vessel is marked "Food or Food
Compatible Only" in conformance with rules adopted under RCW 69.04.960.

(3) No person may transport in intrastate commerce a substance in bulk form other than food or a substance on a list adopted under RCW 69.04.960 in the cargo carrying portion of a vehicle or vessel marked "Food or Food Compatible Only."

(4) This section does not apply to the transportation of a raw agricultural commodity from the point of its production to the facility at which the commodity is first processed or packaged. [1990 c 202 § 2.]

69.04.960 Transport of bulk foods—Compatible substances—Cleaning vehicle or vessel—Vehicle or vessel marking. (1) The director of agriculture and the secretary of health shall jointly adopt by rule:

(a) A list of food compatible substances other than food that may be transported in bulk form as cargo in a vehicle or vessel that is also used, on separate occasions, to transport food in bulk form as cargo. The list shall contain those substances that the director and the secretary determine will not pose a health hazard if food in bulk form were transported in the vehicle or vessel after it transported the substance. In making this determination, the director and the secretary shall assume that some residual portion of the substance will remain in the cargo carrying portion of the vehicle or vessel when the food is transported;

(b) The procedures to be used to clean the vehicle or vessel after transporting the substance and prior to transporting the food;

(c) The form of the certificates to be used under RCW 69.04.965; and

(d) Requirements for the "Food or Food Compatible Only" marking which must be borne by a vehicle or vessel under RCW 69.04.955 or 69.04.965.

(2) In developing and adopting rules under this section and RCW 69.04.970, the director and the secretary shall consult with the secretary of transportation, the chief of the state patrol, the chair of the utilities and transportation commission, and representatives of the vehicle and vessel transportation industries, food processors, and agricultural commodity organizations. [1990 c 202 § 3.]

69.04.965 Transport of bulk foods—Transports not constituting violations. Transporting food as cargo in bulk form in intrastate commerce in a vehicle or vessel that has previously been used to transport in bulk form a cargo other than food does not constitute a violation of RCW 69.04.955 if:

(1) The cargo is a food compatible substance contained on the list adopted by the director and secretary under RCW 69.04.960;

(2) The vehicle or vessel has been cleaned as required by the rules adopted under RCW 69.04.960;

(3) The vehicle or vessel is marked "Food or Food Compatible" in conformance with rules adopted under RCW 69.04.960; and

(4) A certificate accompanies the vehicle or vessel when the food is transported by other than railroad car which attests, under penalty of perjury, to the fact that the vehicle or vessel has been cleaned as required by those rules and is dated and signed by the party responsible for that cleaning. Such certificates shall be maintained by the owner of the vehicle or vessel for not less than three years and shall be available for inspection concerning compliance with RCW 69.04.950 through 69.04.980. The director of agriculture and the secretary of health shall jointly adopt rules requiring such certificates for the transportation of food under this section by railroad car and requiring such certificates to be available for inspection concerning compliance with RCW 69.04.950 through 69.04.980. Forms for the certificates shall be provided by the department of agriculture. [1990 c 202 § 4.]

69.04.970 Transport of bulk foods—Substances rendering vehicle or vessel permanently unsuitable for bulk food transport—Procedures to rehabilitate vehicles and vessels. The director of agriculture and the secretary of health shall jointly adopt by rule:

(1) A list of substances which, if transported in bulk form in the cargo carrying portion of a vehicle or vessel, render the vehicle or vessel permanently unsuitable for use in transporting food in bulk form because the prospect that any residue might be present in the vehicle or vessel when it transports food poses a hazard to the public health; and

(2) Procedures to be used to rehabilitate a vehicle or vessel that has been used to transport a substance other than a substance contained on a list adopted under RCW 69.04.960 or under subsection (1) of this section. The procedures shall ensure that transporting food in the cargo carrying portion of the vehicle or vessel after its rehabilitation will not pose a health hazard. [1990 c 202 § 5.]

69.04.975 Transport of bulk foods—Rehabilitation of vehicles and vessels—Inspection—Certification—Marking—Costs. A vehicle or vessel that has been used to transport a substance other than food or a substance contained on the lists adopted by the director and secretary under RCW 69.04.960 and 69.04.970, may be rehabilitated and used to transport food only if:

(1) The vehicle or vessel is rehabilitated in accordance with the procedures established by the director and secretary in RCW 69.04.970;

(2) The vehicle or vessel is inspected by the department of agriculture, and the department determines that transporting food in the cargo carrying portion of the vehicle or vessel will not pose a health hazard;

(3) A certificate accompanies the vehicle or vessel certifying that the vehicle or vessel has been rehabilitated and inspected and is authorized to transport food, and is dated and signed by the director of agriculture, or an authorized agent of the director. Such certificates shall be maintained for the life of the vehicle by the owner of the vehicle or vessel, and shall be available for inspection concerning compliance with RCW 69.04.950 through 69.04.980. Forms for the certificates shall be provided by the department of agriculture; and

(4) The vehicle or vessel is marked as required by RCW 69.04.955 or is marked and satisfies the requirements of RCW 69.04.965 which are not inconsistent with the rehabilitation authorized by this section.

No vehicle or vessel that has transported in bulk form a substance contained on the list adopted under RCW 69.04.970 qualifies for rehabilitation.
Chapter 69.06 RCW

FOOD AND BEVERAGE ESTABLISHMENT WORKERS’ PERMITS

Sections
69.06.010 Food and beverage service worker’s permit—Filing, duration—Minimum training requirements.
69.06.020 Permit exclusive and valid throughout state—Fee.
69.06.030 Diseased persons—May not work—Employer may not hire.
69.06.040 Application of chapter to retail food establishments.
69.06.045 Application of chapter to temporary food service establishments.
69.06.050 Permit to be secured within fourteen days from time of employment.
69.06.060 Penalty.
69.06.070 Limited duty permit.
69.06.080 Chapter not applicable to persons who meet requirements of RCW 70.128.250.

69.06.010 Food and beverage service worker’s permit—Filing, duration—Minimum training requirements.

It shall be unlawful for any person to be employed in the handling of unwrapped or unpackaged food unless he or she shall furnish and place on file with the person in charge of such establishment, a food and beverage service worker’s permit, as prescribed by the state board of health. Such permit shall be kept on file by the employer or kept by the employee on his or her person and open for inspection at all reasonable hours by authorized public health officials. Such permit shall be returned to the employee upon termination of employment. Initial permits, including limited duty permits, shall be valid for two years from the date of issuance. Subsequent renewal permits shall be valid for three years from the date of issuance, except an employee may be granted a renewal permit that is valid for five years from the date of issuance if the employee demonstrates that he or she has obtained additional food safety training prior to renewal of the permit. Rules establishing minimum training requirements must be adopted by the state board of health and developed by the department of health in conjunction with local health jurisdictions and representatives of the food service industry. [1998 c 136 § 1; 1987 c 223 § 5; 1957 c 197 § 1.]

69.06.020 Permit exclusive and valid throughout state—Fee. The permit provided in RCW 69.06.010 or 69.06.070 shall be valid in every city, town and county in the state, for the period for which it is issued, and no other health certificate shall be required of such employees by any municipal corporation or political subdivision of the state. The cost of the permit shall be uniform throughout the state and shall be in that amount set by the state board of health. The cost of the permit shall reflect actual costs of food worker training and education, administration of the program, and testing of applicants. The state board of health shall periodically review the costs associated with the permit program and adjust the fee accordingly. The board shall also ensure that the fee is not set at an amount that would prohibit low-income persons from obtaining permits. [1998 c 136 § 3; 1987 c 223 § 6; 1957 c 197 § 2.]

69.06.030 Diseased persons—May not work—Employer may not hire. It shall be unlawful for any person afflicted with any contagious or infectious disease that may be transmitted by food or beverage to work in or about any place where unwrapped or unpackaged food and/or beverage products are prepared or sold, or offered for sale for human consumption and it shall be unlawful for any person knowingly to employ a person so afflicted. Nothing in this section eliminates any authority or requirement to control or suppress communicable diseases pursuant to chapter 70.05 RCW and RCW 43.20.050(2)(e). [1998 c 136 § 4; 1957 c 197 § 3.]

*Reviser’s note: RCW 43.20.050 was amended by 2009 c 495 § 1, changing subsection (2)(e) to subsection (2)(f).

69.06.040 Application of chapter to retail food establishments. This chapter shall apply to any retail establishment engaged in the business of food handling or food service. [1987 c 223 § 7; 1957 c 197 § 4.]

69.06.045 Application of chapter to temporary food service establishments. As used in this section, “temporary food service establishment” means a food service establishment operating at a fixed location for a period of time of not more than twenty-one consecutive days in conjunction with a single event or celebration. This chapter applies to temporary food service establishments with the following exceptions:

(1) Only the operator or person in charge of a temporary food service establishment shall be required to secure a food and beverage service workers’ permit; and

(2) The operator or person in charge of a temporary food service establishment shall secure a valid food and beverage service workers’ permit before commencing the food handling operation. [1987 c 223 § 8.]

69.06.050 Permit to be secured within fourteen days from time of employment. Individuals under this chapter must obtain a food and beverage service workers’ permit within fourteen days from commencement of employment.
 Individuals under this chapter may work for up to fourteen calendar days without a food and beverage service workers' permit, provided that they receive information or training regarding safe food handling practices from the employer prior to commencement of employment. Documentation that the information or training has been provided to the individual must be kept on file by the employer. [1998 c 136 § 5; 1957 c 197 § 5.]

69.06.060 Penalty. Any violation of the provisions of this chapter shall be a misdemeanor. [1957 c 197 § 6.]

69.06.070 Limited duty permit. The local health officer may issue a limited duty permit when necessary to reasonably accommodate a person with a disability. The limited duty permit must specify the activities that the permit holder may perform, and must include only activities having low public health risk. [1998 c 136 § 2.]

69.06.080 Chapter not applicable to persons who meet requirements of RCW 70.128.250. Except for the food safety training standards adopted by the state board of health under RCW 69.06.010, the provisions of this chapter do not apply to persons who work in adult family homes and successfully complete training and continuing education as required by RCW 70.128.250. [2005 c 505 § 7.]

Chapter 69.07 RCW  
WASHINGTON FOOD PROCESSING ACT

Sections
69.07.005 Legislative declaration.
69.07.010 Definitions.
69.07.020 Enforcement—Rules—Adoption—Contents—Standards.
69.07.040 Food processing license—Waiver if licensed under chapter 15.36 RCW—Expiration date—Application, contents—Fee.
69.07.050 Renewal of license—Additional fee, when.
69.07.060 Denial, suspension, or revocation of license—Grounds.
69.07.065 Suspension of license summarily—Reinstatement.
69.07.070 Rules and regulations, hearings subject to Administrative Procedure Act.
69.07.080 Inspections by department—Access—When.
69.07.085 Sanitary certificates—Fee.
69.07.095 Authority of director and personnel.
69.07.100 Establishments exempted from provisions of chapter.
69.07.103 Poultry—Slaughter, preparation, sale—One thousand or fewer—Special permit—Rules—Fee.
69.07.110 Enforcement of chapter.
69.07.120 Disposition of money into food processing inspection account.
69.07.135 Unlawful to sell or distribute food from unlicensed processor.
69.07.140 Violations—Warning notice.
69.07.150 Violations—Penalties.
69.07.160 Authority of director and department under chapter 69.04 RCW not impaired by any provision of chapter 69.07 RCW.
69.07.170 Definitions.
69.07.180 Bottled water labeling standards.
69.07.190 Bottled soft drinks, soda, or seltzer exempt from bottled water labeling requirements.
69.07.900 Chapter is cumulative and nonexclusive.
69.07.910 Severability—1967 ex.s.c 121.
69.07.920 Short title.

69.07.005 Legislative declaration. The processing of food intended for public consumption is important and vital to the health and welfare both immediate and future and is hereby declared to be a business affected with the public interest. The provisions of this chapter [1991 c 137] are enacted to safeguard the consuming public from unsafe, adulterated, or misbranded food by requiring licensing of all food processing plants as defined in this chapter and setting forth the requirements for such licensing. [1991 c 137 § 1.]

69.07.010 Definitions. For the purposes of this chapter:
(1) "Department" means the department of agriculture of the state of Washington;
(2) "Director" means the director of the department;
(3) "Food" means any substance used for food or drink by any person, including ice, bottled water, and any ingredient used for components of any such substance regardless of the quantity of such component;
(4) "Sale" means selling, offering for sale, holding for sale, preparing for sale, trading, bartering, offering a gift as an inducement for sale of, and advertising for sale in any media;
(5) "Food processing" means the handling or processing of any food in any manner in preparation for sale for human consumption: PROVIDED, That it shall not include fresh fruit or vegetables merely washed or trimmed while being prepared or packaged for sale in their natural state;
(6) "Food processing plant" includes but is not limited to any premises, plant, establishment, building, room, area, facilities and the appurtenances thereto, in whole or in part, where food is prepared, handled or processed in any manner for distribution or sale for resale by retail outlets, restaurants, and any such other facility selling or distributing to the ultimate consumer: PROVIDED, That, as set forth herein, establishments processing foods in any manner for resale shall be considered a food processing plant as to such processing;
(7) "Food service establishment" shall mean any fixed or mobile restaurant, coffee shop, cafeteria, short order cafe, luncheonette, grill, tearoom, sandwich shop, soda fountain, tavern, bar, cocktail lounge, night club, roadside stand, industrial-feeding establishment, retail grocery, retail food market, retail meat market, retail bakery, private, public, or nonprofit organization routinely serving food, catering kitchen, commissary or similar place in which food or drink is prepared for sale or for service on the premises or elsewhere, and any other eating or drinking establishment or operation where food is served or provided for the public with or without charge.

For the purpose of this chapter any custom cannery or processing plant where raw food products, food, or food products are processed for the owner thereof, or the food processing facilities are made available to the owners or persons in control of raw food products or food or food products for processing in any manner, shall be considered to be food processing plants;
(8) "Person" means an individual, partnership, corporation, or association. [1992 c 34 § 3; 1991 c 137 § 2; 1967 ex.s.c 121 § 1.]

Additional notes found at www.leg.wa.gov

69.07.020 Enforcement—Rules—Adoption—Contents—Standards. (1) The department shall enforce and carry out the provisions of this chapter, and may adopt the necessary rules to carry out its purposes.
(2) Such rules may include:
(a) Standards for temperature controls in the storage of foods, so as to provide proper refrigeration.
(b) Standards for temperatures at which low acid foods must be processed and the length of time such temperatures must be applied and at what pressure in the processing of such low acid foods.

(c) Standards and types of recording devices that must be used in providing records of the processing of low acid foods, and how they shall be made available to the department of agriculture for inspection.

(d) Requirements for the keeping of records of the temperatures, times and pressures at which foods were processed, or for the temperatures at which refrigerated products were stored by the licensee and the furnishing of such records to the department.

(e) Standards that must be used to establish the temperature and purity of water used in the processing of foods.

[1969 c 68 § 1; 1967 ex.s. c 121 § 2.]

69.07.040 Food processing license—Waiver if licensed under chapter 15.36 RCW—Expiration date—Application, contents—Fee. It shall be unlawful for any person to operate a food processing plant or process foods in the state without first having obtained an annual license from the department, which shall expire on a date set by rule by the director. License fees shall be prorated where necessary to accommodate staggering of expiration dates. Application for a license shall be on a form prescribed by the director and accompanied by the license fee. The license fee is determined by computing the gross annual sales for the accounting year immediately preceding the license year. If the license is for a new operator, the license fee shall be based on an estimated gross annual sales for the initial license period.

If gross annual sales are: The license fee is:

<table>
<thead>
<tr>
<th>Sales Range</th>
<th>Fee</th>
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<tbody>
<tr>
<td>$0 to $50,000</td>
<td>$55.00</td>
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<tr>
<td>$50,001 to $500,000</td>
<td>$110.00</td>
</tr>
<tr>
<td>$500,001 to $1,000,000</td>
<td>$220.00</td>
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<tr>
<td>$1,000,001 to $5,000,000</td>
<td>$385.00</td>
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<tr>
<td>$5,000,001 to $10,000,000</td>
<td>$550.00</td>
</tr>
<tr>
<td>Greater than $10,000,000</td>
<td>$825.00</td>
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Such application shall include the full name of the applicant for the license and the location of the food processing plant he or she intends to operate. If such applicant is an individual, receiver, trustee, firm, partnership, association or corporation, the full name of each member of the firm or partnership, or names of the officers of the association or corporation shall be given on the application. Such application shall further state the principal business address of the applicant in the state and elsewhere and the name of a person domiciled in this state authorized to receive and accept service of summons of legal notices of all kinds for the applicant. The application shall also specify the type of food to be processed and the method or nature of processing operation or preservation of that food and any other necessary information. Upon the approval of the application by the director and compliance with the provisions of this chapter, including the applicable regulations adopted hereunder by the department, the applicant shall be issued a license or renewal thereof.

Licenses shall be issued to cover only those products, processes, and operations specified in the license application and approved for licensing. Wherever a license holder wishes to engage in processing a type of food product that is different than the type specified on the application supporting the licensee’s existing license and processing that type of food product would require a major addition to or modification of the licensee’s processing facilities or has a high potential for harm, the licensee shall submit an amendment to the current license application. In such a case, the licensee may engage in processing the new type of food product only after the amendment has been approved by the department.

If upon investigation by the director, it is determined that a person is processing food for retail sale and is not under permit, license, or inspection by a local health authority, then that person may be considered a food processor and subject to the provisions of this chapter. The director may waive the licensure requirements of this chapter for a person’s operations at a facility if the person has obtained a milk processing plant license under chapter 15.36 RCW to conduct the same or a similar operation at the facility. [1995 c 374 § 21. Prior: 1993 sp.s. c 19 § 11; 1993 c 212 § 2; 1992 c 160 § 3; 1991 c 137 § 3; 1988 c 5 § 1; 1969 c 68 § 2; 1967 ex.s. c 121 § 4.]

Additional notes found at www.leg.wa.gov

69.07.050 Renewal of license—Additional fee, when. If the application for renewal of any license provided for under this chapter is not filed prior to the expiration date as established by rule by the director, an additional fee of ten percent of the cost of the license shall be assessed and added to the original fee and shall be paid by the applicant before the renewal license shall be issued: PROVIDED, That such additional fee shall not be charged if the applicant furnishes an affidavit certifying that he or she has not operated a food processing plant or processed foods subsequent to the expiration of his or her license. [1992 c 160 § 4; 1991 c 137 § 4; 1988 c 5 § 2; 1967 ex.s. c 121 § 5.]

69.07.060 Denial, suspension, or revocation of license—Grounds. The director may, subsequent to a hearing thereon, deny, suspend, or revoke any license provided for in this chapter if he or she determines that an applicant has committed any of the following acts:

(1) Refused, neglected, or failed to comply with the provisions of this chapter, the rules and regulations adopted hereunder, or any lawful order of the director.

(2) Refused, neglected, or failed to keep and maintain records required by this chapter, or to make such records available when requested pursuant to the provisions of this chapter.

(3) Refused the department access to any portion or area of the food processing plant for the purpose of carrying out the provisions of this chapter.

(4) Refused the department access to any records required to be kept under the provisions of this chapter.

(5) Refused, neglected, or failed to comply with any provisions of chapter 69.04 RCW, Washington food, drug, and cosmetic act, or any regulations adopted thereunder.

The provisions of this section requiring that a hearing be conducted before an action may be taken against a license do not apply to an action taken under RCW 69.07.065. [2012 c 117 § 345; 1991 c 137 § 5; 1979 c 154 § 19; 1967 ex.s. c 121 § 6.]

Additional notes found at www.leg.wa.gov
69.07.065 Suspension of license summarily—Reinstatement. (1) Whenever the director finds an establishment operating under conditions that constitute an immediate danger to public health or whenever the licensee or any employee of the licensee actively prevents the director or the director’s representative, during an onsite inspection, from determining whether such a condition exists, the director may summarily suspend, pending a hearing, a license provided for in this chapter.

(2) Whenever a license is summarily suspended, the holder of the license shall be notified in writing that the license is, upon service of the notice, immediately suspended and that prompt opportunity for a hearing will be provided.

(3) Whenever a license is summarily suspended, food processing operations shall immediately cease. However, the director may reinstate the license when the condition that caused the suspension has been abated to the director’s satisfaction. [1991 c 137 § 6.]

69.07.070 Rules and regulations, hearings subject to Administrative Procedure Act. The adoption of any rules and regulations under the provisions of this chapter, or the holding of a hearing in regard to a license issued or which may be issued under the provisions of this chapter shall be subject to the applicable provisions of chapter 34.05 RCW, the Administrative Procedure Act, as enacted or hereafter amended. [1967 ex.s. c 121 § 7.]

69.07.080 Inspections by department—Access—When. For purpose of determining whether the rules adopted pursuant to RCW 69.07.020, as now or hereafter amended are complied with, the department shall have access for inspection purposes to any part, portion or area of a food processing plant, and any records required to be kept under the provisions of this chapter or rules and regulations adopted hereunder. Such inspection shall, when possible, be made during regular business hours or during any working shift of said food processing plant. The department may, however, inspect such food processing plant at any time when it has received information that an emergency affecting the public health has arisen and such food processing plant is or may be involved in the matters causing such emergency. [1969 c 68 § 3; 1967 ex.s. c 121 § 8.]

69.07.085 Sanitary certificates—Fee. The department may issue sanitary certificates to food processors under this chapter subject to such requirements as it may establish by rule. The fee for issuance shall be fifty dollars per certificate. Fees collected under this section shall be deposited in the agricultural local fund. [1995 c 374 § 23; 1988 c 254 § 9.]

Additional notes found at www.leg.wa.gov

69.07.095 Authority of director and personnel. The director or the director’s deputies, assistants, and inspectors are authorized to do all acts and things necessary to carry out the provisions of this chapter, including the taking of verified statements. The department personnel are empowered to administer oaths of verification on the statement. [1991 c 137 § 7.]

69.07.100 Establishments exempted from provisions of chapter. (1) The provisions of this chapter shall not apply to establishments issued a permit or licensed under the provisions of:

(a) Chapter 69.25 RCW, the Washington wholesome eggs and egg products act;
(b) Chapter 69.28 RCW, the Washington state honey act;
(c) Chapter 16.49 RCW, the meat inspection act;
(d) Chapter 77.65 RCW, relating to the direct retail endorsement for wild-caught seafood;
(e) Chapter 69.22 RCW, relating to cottage food operations;
(f) Title 66 RCW, relating to alcoholic beverage control; and
(g) Chapter 69.30 RCW, the sanitary control of shellfish act.

(2) If any such establishments process foods not specifically provided for in the above entitled acts, the establishments are subject to the provisions of this chapter.

(3) The provisions of this chapter do not apply to restaurants or food service establishments. [2011 c 281 § 13; 2002 c 301 § 10; 1995 c 374 § 22; 1988 c 5 § 4; 1983 c 3 § 168; 1967 ex.s. c 121 § 10.]

Finding—Effective date—2002 c 301: See notes following RCW 77.65.510.

Additional notes found at www.leg.wa.gov

69.07.103 Poultry—Slaughter, preparation, sale—One thousand or fewer—Special permit—Rules—Fee. (1) A special permit issued by the department under this section is required for the slaughter, preparation, and sale of one thousand or fewer poultry in a calendar year by a poultry producer for the sale of whole raw poultry directly to the ultimate consumer at the producer’s farm. Activities conducted under the permit are exempt from any other licensing requirements of this chapter.

(2)(a) The department must adopt by rule requirements for the permit. The requirements must be generally patterned after those established by the state board of health for temporary food service establishments, but must be tailored specifically to poultry slaughter, preparation, and sale activities. The requirements must include, but are not limited to, those for: Cooling procedures, when applicable; sanitary facilities, equipment, and utensils; clean water; washing and other hygienic practices; and waste and wastewater disposal.

(b) A permit expires December 31st and may be issued for either one or two years as requested by the permit applicant upon payment of the applicable fee in accordance with subsection (4) of this section.

(3) The department shall conduct such inspections as are reasonably necessary to ensure compliance with permit requirements.

(4) The fee for a special permit is seventy-five dollars for one year, or one hundred twenty-five dollars for two years. [2009 c 114 § 1; 2003 c 397 § 2.]

69.07.110 Enforcement of chapter. The department may use all the civil remedies provided for in chapter 69.04 RCW (The Uniform Washington Food, Drug, and Cosmetic Act) in carrying out and enforcing the provisions of this chapter. [1967 ex.s. c 121 § 11.]
69.07.120 Disposition of money into food processing inspection account. All moneys received by the department under the provisions of this chapter and chapter 69.22 RCW shall be paid into the food processing inspection account hereby created within the agricultural local fund established in RCW 43.23.230 and shall be used solely to carry out the provisions of this chapter and chapters 69.22 and 69.04 RCW. [2011 c 281 § 12; 1992 c 160 § 5; 1967 ex.s.c 121 § 12.]

69.07.135 Unlawful to sell or distribute food from unlicensed processor. It shall be unlawful to resell, to offer for resale, or to distribute for resale in intrastate commerce any food processed in a food processing plant, which has not obtained a license, as provided for in this chapter, once notification by the director has been given to the person or persons reselling, offering, or distributing food for resale, that said food is from an unlicensed processing operation. [1991 c 137 § 8.]

69.07.140 Violations—Warning notice. Nothing in this chapter shall be construed as requiring the department to report for prosecution violations of this chapter when it believes that the public interest will best be served by a suitable notice of warning in writing. [1967 ex.s.c 121 § 14.]

69.07.150 Violations—Penalties. (1)(a) Except as provided in (b) of this subsection, any person violating any provision of this chapter or any rule or regulation adopted hereunder is guilty of a misdemeanor.

(b) A second or subsequent violation is a gross misdemeanor. Any offense committed more than five years after a previous conviction shall be considered a first offense.

(2) Whenever the director finds that a person has committed a violation of any of the provisions of this chapter, and that violation has not been punished pursuant to subsection (1) of this section, the director may impose upon and collect from the violator a civil penalty not exceeding one thousand dollars per violation. Each violation shall be a separate and distinct offense. [2003 c 53 § 316; 1991 c 137 § 9; 1967 ex.s.c 121 § 15.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.07.160 Authority of director and department under chapter 69.04 RCW not impaired by any provision of chapter 69.07 RCW. The authority granted to the director and to the department under the provisions of the Uniform Washington Food, Drug, and Cosmetic Act (chapter 69.04 RCW), as now or hereafter amended, shall not be deemed to be reduced or otherwise impaired as a result of any provision or provisions of the Washington Food Processing Act (chapter 69.07 RCW). [1969 c 68 § 4.]

69.07.170 Definitions. As used in RCW 69.07.180 and 69.07.190:

(1) "Artesian water" means bottled water from a well tapping a confined aquifer in which the water level stands above the water table. "Artesian water" shall meet the requirements of "natural water."

(2) "Bottled water" means water that is placed in a sealed container or package and is offered for sale for human consumption or other consumer uses.

(3) "Carbonated water" or "sparkling water" means bottled water containing carbon dioxide.

(4) "Department" means the department of agriculture.

(5) "Distilled water" means bottled water that has been produced by a process of distillation and meets the definition of purified water in the most recent edition of the United States Pharmacopeia.

(6) "Drinking water" means bottled water obtained from an approved source that has at minimum undergone treatment consisting of filtration, activated carbon or particulate, and ozonization or an equivalent disinfection process, or that meets the requirements of the federal safe drinking water act of 1974 as amended and complies with all department of health rules regarding drinking water.

(7) "Mineral water" means bottled water that contains not less than five hundred parts per million total dissolved solids. "Natural mineral water" shall meet the requirements of "natural water."

(8) "Natural water" means bottled spring, mineral, artesian, or well water that is derived from an underground formation and may be derived from a public water system as defined in RCW 70.119A.020 only if that supply has a single source such as an actual spring, artesian well, or pumped well, and has not undergone any treatment that changes its original chemical makeup except ozonization or an equivalent disinfection process.

(9) "Plant operator" means a person who owns or operates a bottled water plant.

(10) "Purified water" means bottled water produced by distillation, deionization, reverse osmosis, or other suitable process and that meets the definition of purified water in the most recent edition of the United States Pharmacopeia. Water that meets this definition and is vaporized, then condensed, may be labeled "distilled water."

(11) "Spring water" means water derived from an underground formation from which water flows naturally to the surface of the earth. "Spring water" shall meet the requirements of "natural water."

(12) "Water dealer" means a person who imports bottled water or causes bulk water to be transported for bottling for human consumption or other consumer uses.

(13) "Well water" means water from a hole bored, drilled, or otherwise constructed in the ground that taps the water of an aquifer. "Well water" shall meet the requirements of "natural water." [1992 c 34 § 1.]

Additional notes found at www.leg.wa.gov

69.07.180 Bottled water labeling standards. All bottled water must conform to applicable federal and state labeling laws and be labeled in compliance with the following standards:

(1) Mineral water may be labeled "mineral water." Bottled water to which minerals are added shall be labeled so as to disclose that minerals are added, and may not be labeled "natural mineral water."

(2) Spring water may be labeled "spring water" or "natural spring water."

[Title 69 RCW—page 24]
Food Storage Warehouses

Chapter 69.10 RCW

FOOD STORAGE WAREHOUSES

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69.10.020 Exemption from licensure—Independent inspection—Report to department.
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69.10.035 Immediate danger to public health—Summary suspending license—Written notification—Hearing—Reinstatement of license.
69.10.040 Unlicensed food storage warehouse—Unlawful to sell, offer for sale, or distribute in intrastate commerce.
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69.10.050 Civil remedies—Restrictions on civil penalties—Fee limitations for inspections and analyses.
69.10.055 Rules.
69.10.060 Director and deputies, assistants, and inspectors authorized to act—May take verified statements.

69.10.005 Definitions. For the purpose of this chapter:

(1) "Food storage warehouse" means any premises, establishment, building, room area, facility, or place, in whole or in part, where food is stored, kept, or held for wholesale distribution to other wholesalers or to retail outlets, restaurants, and any such other facility selling or distributing to the ultimate consumer. Food storage warehouses include, but are not limited to, facilities where food is kept or held refrigerated or frozen and include facilities where food is stored to the account of another firm and/or is owned by the food storage warehouse. "Food storage warehouse" does not include grain elevators or fruit and vegetable storage and packing houses that store, pack, and ship fresh fruit and vegetables even though they may use refrigerated or controlled atmosphere storage practices in their operation. However, this chapter applies to multiple food storage operations that also distribute or ripen fruits and vegetables.

(2) "Department" means the Washington department of agriculture.

(3) "Director" means the director of the Washington department of agriculture.

(4) "Food" means the same as defined in RCW 69.04.008.

(5) "Independent sanitation consultant" means an individual, partnership, cooperative, or corporation that by reason of education, certification, and experience has satisfactorily demonstrated expertise in food and dairy sanitation and is approved by the director to advise on such areas including, but not limited to: Principles of cleaning and sanitizing food processing plants and equipment; rodent, insect, bird, and other pest control; principals [principles] of hazard analysis critical control point; basic food product labeling; principles of proper food storage and protection; proper personnel work practices and attire; sanitary design, construction, and installation of food plant facilities, equipment, and utensils; and other pertinent food safety issues. [1995 c 374 § 8.]

69.10.010 Inspection of food storage warehouses—Powers of director. The director or his or her representative may inspect food storage warehouses for compliance with the
provisions of chapter 69.04 RCW and the rules adopted under chapter 69.04 RCW as deemed necessary by the director. Any food storage warehouse found to not be in substantial compliance with chapter 69.04 RCW and the rules adopted under chapter 69.04 RCW will be reinspected as deemed necessary by the director to determine compliance. This does not preclude the director from using any other remedies as provided under chapter 69.04 RCW to gain compliance or to embargo products as provided under RCW 69.04.110 to protect the public from adulterated foods. [1995 c 374 § 9.]

69.10.015 Annual license required—Director's duties—Fee—Application—Renewal. Except as provided in this section and RCW 69.10.020, it shall be unlawful for any person to operate a food storage warehouse in the state without first having obtained an annual license from the department, which shall expire on a date set by rule by the director. Application for a license or license renewal shall be on a form prescribed by the director and accompanied by the license fee. The license fee is fifty dollars.

For a food storage warehouse that has been inspected on at least an annual basis for compliance with the provisions of the current good manufacturing practices (Title 21 C.F.R. part 110) by a federal agency or by a state agency acting on behalf of and under contract with a federal agency and that is not exempted from licensure by RCW 69.10.020, the annual license fee for the warehouse is twenty-five dollars.

The application shall include the full name of the applicant for the license and the location of the food storage warehouse he or she intends to operate. If such applicant is an individual, receiver, trustee, firm, partnership, association, or corporation, the full name of each member of the firm or partnership, or names of the officers of the association or corporation must be given on the application. The application shall further state the principal business address of the applicant in the state and elsewhere and the name of a person domiciled in this state authorized to receive and accept service of summons of legal notices of all kinds for the applicant. Upon the approval of the application by the director and compliance with the provisions of this chapter, including the applicable regulations adopted under this chapter by the department, the applicant shall be issued a license or renewal thereof. The director shall waive licensure under this chapter for firms that are licensed under the provisions of chapter 69.07 or 15.36 RCW. [1995 c 374 § 10.]

69.10.020 Exemption from licensure—Independent inspection—Report to department. A food storage warehouse that is inspected for compliance with the current good manufacturing practices (Title 21 C.F.R. part 110) on at least an annual basis by an independent sanitation consultant approved by the department shall be exempted from licensure under this chapter.

A report identifying the inspector and the inspecting entity, the date of the inspection, and any violations noted on such inspection shall be forwarded to the department by the food storage warehouse within sixty days of the completion of the inspection. An inspection shall be conducted and an inspection report for a food storage warehouse shall be filed with the department at least once every twelve months or the warehouse shall be licensed under this chapter and inspected by the department for a period of two years. [1995 c 374 § 11.]

69.10.025 Application for renewal of license after expiration date—Additional fee. If the application for renewal of any license provided for under this chapter is not filed prior to the expiration date as established by rule by the director, an additional fee of ten percent of the cost of the license shall be assessed and added to the original fee and must be paid by the applicant before the renewal license is issued. [1995 c 374 § 12.]

69.10.030 Director may deny, suspend, or revoke license—Actions by applicant—Hearing required. The director may, subsequent to a hearing thereon, deny, suspend, or revoke any license provided for in this chapter if he or she determines that an applicant has committed any of the following acts:

1. Refused, neglected, or failed to comply with the provisions of this chapter, the rules adopted under this chapter, or any lawful order of the director;
2. Refused, neglected, or failed to keep and maintain records required by this chapter, or to make such records available if requested pursuant to the provisions of this chapter;
3. Refused the department access to any portion or area of the food storage warehouse for the purpose of carrying out the provisions of this chapter;
4. Refused the department access to any records required to be kept under the provisions of this chapter;
5. Refused, neglected, or failed to comply with any provisions of chapter 69.04 RCW, Washington food, drug, and cosmetic act, or any rules adopted under chapter 69.04 RCW.

The provisions of this section requiring that a hearing be conducted before an action may be taken against a license do not apply to an action taken under RCW 69.10.035. [1995 c 374 § 13.]

69.10.035 Immediate danger to public health—Summarily suspending license—Written notification—Hearing—Reinstatement of license. (1) Whenever the director finds a food storage warehouse operating under conditions that constitute an immediate danger to public health or whenever the licensee or any employee of the licensee actively prevents the director or the director’s representative, during an on-site inspection, from determining whether such a condition exists, the director may summarily suspend, pending a hearing, a license provided for in this chapter.

2. Whenever a license is summarily suspended, the holder of the license shall be notified in writing that the license is, upon service of the notice, immediately suspended.

3. Whenever a license is summarily suspended, food distribution operations shall immediately cease. However, the director may reinstate the license if the condition that caused the suspension has been abated to the director’s satisfaction. [1995 c 374 § 14.]

69.10.040 Unlicensed food storage warehouse—Unlawful to sell, offer for sale, or distribute in intrastate commerce. It is unlawful to sell, offer for sale, or distribute...
in intrastate commerce food from or stored in a food storage warehouse that is required to be licensed under this chapter but that has not obtained a license, once notification by the director has been given to the persons selling, offering, or distributing food for sale, that the food is in or from such an unlicensed food storage warehouse. [1995 c 374 § 15.]

69.10.045 Disposition of moneys received under this chapter. All moneys received by the department under provisions of this chapter, except moneys collected for civil penalties levied under this chapter, shall be paid into an account created in the agricultural local fund established in RCW 43.23.230 and shall be used solely to carry out provisions of this chapter and chapter 69.04 RCW. All moneys collected for civil penalties levied under this chapter shall be deposited in the state general fund. [1995 c 374 § 16.]

69.10.050 Civil remedies—Restrictions on civil penalties—Fee limitations for inspections and analyses. (1) Except as provided in subsection (2) of this section, the department may use all the civil remedies provided under chapter 69.04 RCW in carrying out and enforcing the provisions of this chapter.

(2) Civil penalties are intended to be used to obtain compliance and shall not be collected if a warehouse successfully completes a mutually agreed upon compliance agreement with the department. A warehouse that enters into a compliance agreement with the department shall pay only for inspections conducted by the department and any laboratory analyses as required by the inspections as outlined and agreed to in the compliance agreement. In no event shall the fee for these inspections and analyses exceed four hundred dollars per inspection or one thousand dollars in total. [1995 c 374 § 17.]

69.10.055 Rules. (1) The department shall enforce and carry out the provisions of this chapter and may adopt the necessary rules to carry out its purpose.

(2) The adoption of rules under the provisions of this chapter are subject to the applicable provisions of chapter 34.05 RCW, the administrative procedure act. [1995 c 374 § 18.]

69.10.060 Director and deputies, assistants, and inspectors authorized to act—May take verified statements. The director or director’s deputies, assistants, and inspectors are authorized to do all acts and things necessary to carry out the provisions of this chapter, including the taking of verified statements. The department personnel are empowered to administer oaths of verification on the statement. [1995 c 374 § 19.]


Chapter 69.22 RCW

COTTAGE FOOD OPERATIONS

Sections

69.22.010 Definitions.
69.22.020 Requirements—Authority of director.
69.22.030 Permits, permit renewals.
69.22.040 Basic hygiene inspections.
69.22.050 Annual gross sales—Department to determine annual amount.
69.22.060 Access to permitted areas of domestic residence housing cottage food operations—Authority of director.
69.22.070 Cottage foods operations permit—Denial, suspension, or revocation.
69.22.080 Application of administrative procedure act.
69.22.090 Penalties.
69.22.100 Exemption—Provisions of chapter 69.07 RCW or permitting and inspection by local health jurisdiction.
69.22.110 Application of other state or federal laws or local unit of government ordinances not affected.

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

(1) "Cottage food operation" means a person who produces cottage food products only in the home kitchen of that person’s primary domestic residence in Washington and only for sale directly to the consumer.

(2) "Cottage food products" means nonpotentially hazardous baked goods; jams, jellies, preserves, and fruit butters as defined in 21 C.F.R. Sec. 150 as it existed on July 22, 2011; and other nonpotentially hazardous foods identified by the director in rule.

(3) "Department" means the department of agriculture.

(4) "Director" means the director of the department.

(5) "Domestic residence" means a single-family dwelling or an area within a rental unit where a single person or family actually resides. Domestic residence does not include:

(a) A group or communal residential setting within any type of structure; or

(b) An outbuilding, shed, barn, or other similar structure.

(6) "Home kitchen" means a kitchen primarily intended for use by the residents of a home. It may contain one or more stoves or ovens, which may be a double oven, designed for residential use.

(7) "Permitted area" means the portion of a domestic residence housing a home kitchen where the preparation, packaging, storage, or handling of cottage food products occurs.

(8) "Potentially hazardous food" means foods requiring temperature control for safety because they are capable of supporting the rapid growth of pathogenic or toxigenic microorganisms, or the growth and toxin production of Clostridium botulinum. [2011 c 281 § 1.]

69.22.020 Requirements—Authority of director. (1) The director may adopt, by rule, requirements for cottage food operations. These requirements may include, but are not limited to:

(a) The application and renewal of permits under RCW 69.22.030;

(b) Inspections as provided under RCW 69.22.040;

(c) Sanitary procedures;

(d) Facility, equipment, and utensil requirements;

(e) Labeling specificity beyond the requirements of this section;

(f) Requirements for clean water sources and waste and wastewater disposal; and

(g) Requirements for washing and other hygienic practices.

(2) A cottage food operation must package and properly label for sale to the consumer any food it produces, and the...
food may not be repackaged, sold, or used as an ingredient in other foods by a food processing plant, or sold by a food service establishment.

(3) A cottage food operation must place on the label of any food it produces or packages, at a minimum, the following information:
   (a) The name and address of the business of the cottage food operation;
   (b) The name of the cottage food product;
   (c) The ingredients of the cottage food product, in descending order of predominance by weight;
   (d) The net weight or net volume of the cottage food product;
   (e) Allergen labeling as specified by the director in rule;
   (f) If any nutritional claim is made, appropriate labeling as specified by the director in rule;
   (g) The following statement printed in at least the equivalent of eleven-point font size in a color that provides a clear contrast to the background: "Made in a home kitchen that has not been subject to standard inspection criteria."

(4) Cottage food products may only be sold directly to the consumer and may not be sold by internet, mail order, or for retail sale outside the state.

(5) Cottage food products must be stored only in the primary domestic residence. [2011 c 281 § 2.]

69.22.030 Permits, permit renewals. (1) All cottage food operations must be permitted annually by the department on forms developed by the department. All permits and permit renewals must be made on forms developed by the director and be accompanied by an inspection fee as provided in RCW 69.22.040, a seventy-five dollar public health review fee, and a thirty dollar processing fee. All fees must be deposited into the food processing inspection account created in RCW 69.07.120.

(2) In addition to the provision of any information required by the director on forms developed under subsection (1) of this section and the payment of all fees, an applicant for a permit or a permit renewal as a cottage food operation must also provide documentation that all individuals to be involved in the preparation of cottage foods [cottage food products] have secured a food and beverage service worker’s permit under chapter 69.06 RCW.

(3) All cottage food operations permitted under this section must include a signed document attesting, by opting to become permitted, that the permitted cottage food operation expressly grants to the director the right to enter the domestic residence housing the cottage food operation during normal business hours, or at other reasonable times, for the purposes of inspections under this chapter. [2011 c 281 § 3.]

69.22.040 Basic hygiene inspections. (1) The permitted area of all cottage food operations must be inspected for basic hygiene by the director both before initial permitting under RCW 69.22.030 and annually after initial permitting. In addition, the director may inspect the permitted area of a cottage food operation at any time in response to a foodborne outbreak or other public health emergency.

(2) When conducting an annual basic hygiene inspection, the director shall, at a minimum, inspect for the following:

(a) That the permitted cottage food operator understands that no person other than the permittee, or a person under the direct supervision of the permittee, may be engaged in the processing, preparing, packaging, or handling of any cottage food products or be in the home kitchen during the preparation, packaging, or handling of any cottage food products;

(b) That no cottage food preparation, packaging, or handling is occurring in the home kitchen concurrent with any other domestic activities such as family meal preparation, dishwashing, clothes washing or ironing, kitchen cleaning, or guest entertainment;

(c) That no infants, small children, or pets are in the home kitchen during the preparation, packaging, or handling of any cottage food products;

(d) That all food contact surfaces, equipment, and utensils used for the preparation, packaging, or handling of any cottage food products are washed, rinsed, and sanitized before each use;

(e) That all food preparation and food and equipment storage areas are maintained free of rodents and insects; and

(f) That all persons involved in the preparation and packaging of cottage food products:

   (i) Have obtained a food and beverage service worker’s permit under chapter 69.06 RCW;
   (ii) Are not going to work in the home kitchen when ill;
   (iii) Wash their hands before any food preparation and food packaging activities; and
   (iv) Avoid bare hand contact with ready-to-eat foods through the use of single-service gloves, bakery papers, tongs, or other utensils.

(3) The department shall charge an inspection fee of one hundred twenty-five dollars for any initial or annual basic hygiene inspection, which must be deposited into the food processing inspection account created in RCW 69.07.120. An additional inspection fee must be collected for each visit to a cottage food operation for the purposes of conducting an inspection for compliance.

(4) The director may contract with local health jurisdictions to conduct the inspections required under this section. [2011 c 281 § 4.]

69.22.050 Annual gross sales—Department to determine annual amount. (1) The gross sales of cottage food products may not exceed an annual amount set by the department. The determination of the maximum annual gross sales must be computed on the basis of the amount of gross sales within or at a particular domestic residence and may not be computed on a per person basis within or at an individual domestic residence.

(2) If gross sales exceed the maximum annual gross sales amount, the cottage food operation must either obtain a food processing plant license under chapter 69.07 RCW or cease operations.

(3) A cottage food operation exceeding the maximum annual gross sales amount is not entitled to a full or partial refund of any fees paid under RCW 69.22.030 or 69.22.040.

(4) The maximum annual gross sales amount must be established in rule by the department consistent with this subsection. The amount must be set at fifteen thousand dollars until December 31, 2012. Beginning January 1, 2013, the department must increase the fifteen thousand dollar annual
gross sales limit biennially to reflect inflation. The department may determine inflation-based increases in any matter it deems most efficient.

(5) The director may request in writing documentation to verify the annual gross sales figure. [2011 c 281 § 5.]

69.22.060 Access to permitted areas of domestic residence housing cottage food operations—Authority of director. (1) For the purpose of determining compliance with this chapter, the director may access, for inspection purposes, the permitted area of a domestic residence housing a cottage food operation permitted by the director under this chapter. This authority includes the authority to inspect any records required to be kept under the provisions of this chapter.

(2) All inspections must be made at reasonable times and, when possible, during regular business hours.

(3) Should the director be denied access to the permitted area of a domestic residence housing a cottage food operation where access was sought for the purposes of enforcing or administering this chapter, the director may apply to any court of competent jurisdiction for a search warrant authorizing access to the permitted area of a domestic residence housing a permitted cottage food operation, upon which the court may issue a search warrant for the purposes requested.

(4) Any access under this section must be limited to the permitted area and further limited to the purpose of enforcing or administering this chapter. [2011 c 281 § 6.]

69.22.070 Cottage foods operations permit—Denial, suspension, or revocation. (1) After conducting a hearing, the director may deny, suspend, or revoke any permit provided for in this chapter if it is determined that a permittee has committed any of the following acts:

(a) Refused, neglected, or failed to comply with the provisions of this chapter, any rules adopted to administer this chapter, or any lawful order of the director;

(b) Refused, neglected, or failed to keep and maintain records required by this chapter, or to make the records available when requested pursuant to the provisions of this chapter;

(c) Consistent with RCW 69.22.060, refused the director access to the permitted area of a domestic residence housing a cottage food operation for the purpose of carrying out the provisions of this chapter;

(d) Consistent with RCW 69.22.060, refused the department access to any records required to be kept under the provisions of this chapter;

(e) Exceeded the annual income limits provided in RCW 69.22.050.

(2) The director may summarily suspend a permit issued under this chapter if the director finds that a cottage food operation is operating under conditions that constitute an immediate danger to public health or if the director is denied access to the permitted area of a domestic residence housing a cottage food operation and records where the access was sought for the purposes of enforcing or administering this chapter. [2011 c 281 § 7.]

69.22.080 Application of administrative procedure act. The rights, remedies, and procedures respecting the administration of this chapter, including rule making, emergency actions, and permit suspension, revocation, or denial are governed by chapter 34.05 RCW. [2011 c 281 § 8.]

69.22.090 Penalties. (1)(a) Any person engaging in a cottage food operation without a valid permit issued under RCW 69.22.030 or otherwise violating any provision of this chapter, or any rule adopted under this chapter, is guilty of a misdemeanor.

(b) A second or subsequent violation is a gross misdemeanor. Any offense committed more than five years after a previous conviction shall be considered a first offense.

(2) Whenever the director finds that a person has committed a violation of any of the provisions of this chapter, and that violation has not been punished pursuant to subsection (1) of this section, the director may impose upon and collect from the violator a civil penalty not exceeding one thousand dollars per violation per day. Each violation shall be a separate and distinct offense. [2011 c 281 § 9.]

69.22.100 Exemption—Provisions of chapter 69.07 RCW or permitting and inspection by local health jurisdiction. Except as otherwise provided in this chapter, cottage food operations with a valid permit under RCW 69.22.030 are not subject to the provisions of chapter 69.07 RCW or to permitting and inspection by a local health jurisdiction. [2011 c 281 § 10.]

69.22.110 Application of other state or federal laws or local unit of government ordinances not affected. Nothing in this chapter affects the application of any other state or federal laws or any applicable ordinances enacted by any local unit of government. [2011 c 281 § 11.]

Chapter 69.25 RCW

WASHINGTON WHOLESOME EGGS AND EGG PRODUCTS ACT

Sections

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69.25.010 Legislative finding. Eggs and egg products are an important source of the state’s total supply of food, and are used in food in various forms. They are consumed throughout the state and the major portion thereof moves in intrastate commerce. It is essential, in the public interest, that the health and welfare of consumers be protected by the adoption of measures prescribed herein for assuring that eggs and egg products distributed to them and used in products consumed by them are wholesome, otherwise not adulterated, and properly labeled and packaged. Lack of effective regulation for the handling or disposition of unwholesome, otherwise adulterated, or improperly labeled or packaged egg products and certain qualities of eggs is injurious to the public welfare and destroys markets for wholesome, unadulterated, and properly labeled and packaged eggs and egg products and results in sundry losses to producers and processors, as well as injury to consumers. Unwholesome, otherwise adulterated, or improperly labeled or packaged products can be sold at lower prices and compete unfairly with the wholesome, unadulterated, and properly labeled and packaged products, to the detriment of consumers and the public generally. It is hereby found that all egg products and the qualities of eggs which are regulated under this chapter are either in intrastate commerce, or substantially affect such commerce, and that regulation by the director, as contemplated by this chapter, is appropriate to protect the health and welfare of consumers. [1975 1st ex.s. c 201 § 2.]

69.25.020 Definitions. When used in this chapter the following terms shall have the indicated meanings, unless the context otherwise requires:

(1) "Adulterated" applies to any egg or egg product under one or more of the following circumstances:

(a) If it bears or contains any poisonous or deleterious substance which may render it injurious to health; but in case the substance is not an added substance, such article shall not be considered adulterated under this clause if the quantity of such substance in or on such article does not ordinarily render it injurious to health;

(b) If it bears or contains any added poisonous or added deleterious substance (other than one which is: (i) A pesticide chemical in or on a raw agricultural commodity; (ii) a food additive; or (iii) a color additive) which may, in the judgment of the director, make such article unfit for human food;

(c) If it is, in whole or in part, a raw agricultural commodity and such commodity bears or contains a pesticide chemical which is unsafe within the meaning of RCW 69.04.392, as enacted or hereafter amended;

(d) If it bears or contains any food additive which is unsafe within the meaning of RCW 69.04.394, as enacted or hereafter amended;

(e) If it bears or contains any color additive which is unsafe within the meaning of RCW 69.04.396; however, an article which is not otherwise deemed adulterated under subsection (1)(c), (d), or (e) of this section shall nevertheless be deemed adulterated if use of the pesticide chemical, food additive, or color additive, in or on such article, is prohibited by regulations of the director in official plants;

(f) If it consists in whole or in part of any filthy, putrid, or decomposed substance, or if it is otherwise unfit for human food;

(g) If it consists in whole or in part of any damaged egg or eggs to the extent that the egg meat or white is leaking, or it has been contacted by egg meat or white leaking from other eggs;

(h) If it has been prepared, packaged, or held under insanitary conditions whereby it may have become contaminated with filth, or whereby it may have been rendered injurious to health;

(i) If it is an egg which has been subjected to incubation or the product of any egg which has been subjected to incubation;

(j) If its container is composed, in whole or in part, of any poisonous or deleterious substance which may render the contents injurious to health;

(k) If it has been intentionally subjected to radiation, unless the use of the radiation was in conformity with a regulation or exemption in effect pursuant to RCW 69.04.394; or

(l) If any valuable constituent has been in whole or in part omitted or abstracted therefrom; or if any substance has been substituted, wholly or in part therefor; or if damage or inferiority has been concealed in any manner; or if any substance has been added thereto or mixed or packed therewith so as to increase its bulk or weight, or reduce its quality or strength, or make it appear better or of greater value than it is.

(2) "Ambient temperature" means the atmospheric temperature surrounding or encircling shell eggs.

(3) "At retail" means any transaction in intrastate commerce between a retailer and a consumer.

(4) "Candling" means the examination of the interior of eggs by the use of transmitted light used in a partially dark room or place.

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69.25.300 Transfer of moneys in state egg account.

69.25.310 Containers—Marking required—Obliteration of previous markings required for reuse—Temporary use of another handler’s or dealer’s permanent number—Penalty.

69.25.320 Records required, additional—Sales to retailer or food service—Exception—Defense to charged violation—Sale of eggs deteriorated due to storage time—Requirements for storage, display, or transportation.

69.25.900 Savings.

69.25.910 Chapter is cumulative and nonexclusive.

69.25.920 Severability—1975 1st ex.s. c 201.

69.25.930 Short title.
(5) "Capable of use as human food" shall apply to any egg or egg product unless it is denatured, or otherwise identified, as required by regulations prescribed by the director, to deter its use as human food.

(6) "Check" means an egg that has a broken shell or crack in the shell but has its shell membranes intact and contents not leaking.

(7) "Clean and sound shell egg" means any egg whose shell is free of adhering dirt or foreign material and is not cracked or broken.

(8) "Consumer" means any person who purchases eggs for his or her own family use or consumption; or any restaurant, hotel, boarding house, bakery, or other institution or concern which purchases eggs for serving to guests or patrons thereof, or for its own use in cooking or baking.

(9) "Container" or "package" includes any box, can, tin, plastic, or other receptacle, wrapper, or cover.

(10) "Department" means the department of agriculture of the state of Washington.

(11) "Director" means the director of the department or his duly authorized representative.

(12) "Dirty egg" means an egg that has a shell that is unbroken and has adhering dirt or foreign material.

(13) "Egg" means the shell egg of the domesticated chicken, turkey, duck, goose, or guinea, or any other species of fowl.

(14) "Egg handler" or "dealer" means any person who produces, contracts for or obtains possession or control of any eggs or egg products for the purpose of sale to another dealer or retailer, or for processing and sale to a dealer, retailer or consumer. For the purpose of this chapter, "sell" or "sale" includes the following: Offer for sale, expose for sale, have in possession for sale, exchange, barter, trade, or as an inducement for the sale of another product.

(15)(a) "Egg product" means any dried, frozen, or liquid eggs, with or without added ingredients, excepting products which contain eggs only in a relatively small proportion, or historically have not been, in the judgment of the director, considered by consumers as products of the egg food industry, and which may be exempted by the director under such conditions as the director may prescribe to assure that the egg ingredients are not adulterated and are not represented as egg products.

(b) The following products are not included in the definition of "egg product" if they are prepared from eggs or egg products that have been either inspected by the United States department of agriculture or by the department under a cooperative agreement with the United States department of agriculture: Freeze-dried products, imitation egg products, egg substitutes, dietary foods, dried no-bake custard mixes, egg-nog mixes, acidic dressings, noodles, milk and egg dip, cake mixes, French toast, balut and other similar ethnic delicacies, and sandwiches containing eggs or egg products.

(16) "Immediate container" means any consumer package, or any other container in which egg products, not consumer-packaged, are packed.

(17) "Incubator reject" means an egg that has been subjected to incubation and has been removed from incubation during the hatching operations as infertile or otherwise unhatchable.

(18) "Inedible" means eggs of the following descriptions: Black rots, yellow rots, white rots, mixed rots (addled eggs), sour eggs, eggs with green whites, eggs with stuck yolks, moldy eggs, musty eggs, eggs showing blood rings, and eggs containing embryo chicks (at or beyond the blood ring stage).

(19) "Inspection" means the application of such inspection methods and techniques as are deemed necessary by the director to carry out the provisions of this chapter.

(20) "Inspector" means any employee or official of the department authorized to inspect eggs or egg products under the authority of this chapter.

(21) "Intrastate commerce" means any eggs or egg products in intrastate commerce, whether such eggs or egg products are intended for sale, held for sale, offered for sale, sold, stored, transported, or handled in this state in any manner and prepared for eventual distribution in this state, whether at wholesale or retail.

(22) "Leaker" means an egg that has a crack or break in the shell and shell membranes to the extent that the egg contents are exposed or are exuding or free to exude through the shell.

(23) "Loss" means an egg that is unfit for human food because it is smashed or broken so that its contents are leaking; or overheated, frozen, or contaminated; or an incubator reject; or because it contains a bloody white, large meat spots, a large quantity of blood, or other foreign material.

(24) "Master license" means the mechanism established by chapter 19.02 RCW by which master licenses, endorsed for individual state-issued licenses, are issued and renewed utilizing a master application and a master license expiration date common to each renewable license endorsement.

(25) "Misbranded" shall apply to egg products which are not labeled and packaged in accordance with the requirements prescribed by regulations of the director under RCW 69.25.100.

(26) "Official certificate" means any certificate prescribed by regulations of the director for issuance by an inspector or other person performing official functions under this chapter.

(27) "Official device" means any device prescribed or authorized by the director for use in applying any official mark.

(28) "Official inspection legend" means any symbol prescribed by regulations of the director showing that egg products were inspected in accordance with this chapter.

(29) "Official mark" means the official inspection legend or any other symbol prescribed by regulations of the director to identify the status of any article under this chapter.

(30) "Official plant" means any plant which is licensed under the provisions of this chapter, at which inspection of the processing of egg products is maintained by the United States department of agriculture or by the state under cooperative agreements with the United States department of agriculture or by the state.

(31) "Official standards" means the standards of quality, grades, and weight classes for eggs, adopted under the provisions of this chapter.

(32) "Pasteurize" means the subjecting of each particle of egg products to heat or other treatments to destroy harm-
69.25.040 Application of administrative procedure act. The adoption, amendment, modification, or revocation of any rules or regulations under the provisions of this chapter, or the holding of a hearing in regard to a license issued or which may be issued or denied under the provisions of this chapter, shall be subject to the applicable provisions of chapter 34.05 RCW, the administrative procedure act, as now or hereafter amended. [1975 1st ex.s. c 201 § 5.]

69.25.050 Egg handler’s or dealer’s license and number—Branch license—Application, fee, posting required, procedure. (1)(a) No person shall act as an egg handler or dealer without first obtaining an annual license and permanent dealer’s number from the department.

(b) Application for an egg dealer license or egg dealer branch license must be made through the master license system as provided under chapter 19.02 RCW and expires on the master license expiration date. The annual egg dealer license fee is thirty dollars and the annual egg dealer branch license fee is fifteen dollars. A copy of the master license must be posted at each location where the licensee operates. The application must include the full name of the applicant for the license, the location of each facility the applicant intends to operate, and, if applicable, documentation of compliance with RCW 69.25.065 or 69.25.103.

(2) If an applicant is an individual, receiver, trustee, firm, partnership, association or corporation, the full name of each member of the firm or partnership or the names of the officers of the association or corporation shall be given on the application. The application must further state the principal business address of the applicant in the state and elsewhere and the name of a person domiciled in this state authorized to receive and accept service of summons of legal notices of all kinds for the applicant and any other necessary information prescribed by the director.

(3) The applicant must be issued a license or renewal under this section upon the approval of the application and compliance with the provisions of this chapter, including the applicable rules adopted by the department.

(4) The license and permanent egg handler or dealer’s number is nontransferable. [2011 c 306 § 2; 1995 c 374 § 26; 1982 c 182 § 43; 1975 1st ex.s. c 201 § 6.]

Effective date—2011 c 306: See note following RCW 69.25.020.

Master license—Expiration date: RCW 19.02.090.

Master license system definition: RCW 69.25.020(24).

existing licenses or permits registered under, when: RCW 19.02.810.

to include additional licenses: RCW 19.02.110.

Additional notes found at www.leg.wa.gov

69.25.060 Egg handler’s or dealer’s license—Late renewal fee. If the application for the renewal of an egg handler’s or dealer’s license is not filed before the master license expiration date, the master license delinquency fee shall be assessed under chapter 19.02 RCW and shall be paid by the applicant before the renewal license shall be issued. [1982 c 182 § 44; 1975 1st ex.s. c 201 § 7.]

Master license delinquency fee—Rate—Disposition: RCW 19.02.085.

expiration date: RCW 19.02.090.

system—Existing licenses or permits registered under, when: RCW 19.02.810.
69.25.065 Egg handler’s or dealer’s license—Renewal applications—Egg and egg products provided in intrastate commerce produced by commercial egg layer operations—Proof. (1) All new and renewal applications submitted under RCW 69.25.050 before January 1, 2026, must include proof that all eggs and egg products provided in intrastate commerce by the applicant are produced by commercial egg layer operations:

(a) With a current certification under the 2010 version of the United Egg Producers animal husbandry guidelines for United States egg laying flocks for conventional cage systems or cage-free systems or a subsequent version of the guidelines recognized by the department in rule; or

(b) Operated in strict compliance with any standards, adopted by the department in rule, that are equivalent to or more stringent than the standards identified in (a) of this subsection.

(2) All new and renewal applications submitted under RCW 69.25.050 before January 1, 2017, must, in addition to complying with subsection (1) of this section, include proof that all eggs and egg products provided in intrastate commerce by the applicant are produced by commercial egg layer operations whose housing facilities, if built between January 1, 2012, and December 31, 2016, are either:

(a) Approved under, or convertible to, the American humane association facility system plan for enriched colony housing in effect on January 1, 2011, or a subsequent version of the plan recognized by the department in rule and, in addition, are convertible to the standards identified in RCW 69.25.107; or

(b) Operated in strict compliance with any standards, adopted by the department in rule, that are equivalent to or more stringent than the standards identified in (a) of this subsection.

(3) All new and renewal applications submitted under RCW 69.25.050 between January 1, 2017, and December 31, 2025, must, in addition to complying with subsection (1) of this section, include proof that all eggs and egg products provided in intrastate commerce by the applicant are produced by commercial egg layer operations whose housing facilities, if built on or after January 1, 2012, are either:

(a) Approved under the American humane association facility system plan and audit protocol for enriched colony housing in effect on January 1, 2011, or a subsequent version of the plan recognized by the department in rule and, in addition, are operated to the standards identified in RCW 69.25.107; or

(b) Operated in strict compliance with any standards, adopted by the department in rule, that are equivalent to or more stringent than the standards identified in (a) of this subsection.

(4) All new and renewal applications submitted under RCW 69.25.050 on or after January 1, 2026, must include proof that all eggs and egg products provided in intrastate commerce by the applicant are produced by commercial egg layer operations that are either:

(a) Approved under the American humane association facility system plan and audit protocol for enriched colony housing in effect on January 1, 2011, or a subsequent version of the plan recognized by the department in rule and, in addition, are operated to the standards identified in RCW 69.25.107; or

(b) Operated in strict compliance with any standards, adopted by the department in rule, that are equivalent to or more stringent than the standards identified in (a) of this subsection.

(5) The following are exempt from the requirements of subsections (2) and (3) of this section:

(a) Applicants with fewer than three thousand laying chickens; and

(b) Commercial egg layer operations when producing eggs or egg products from turkeys, ducks, geese, guineas, or other species of fowl other than domestic chickens. [2011 c 306 § 3.]

Effective date—2011 c 306: See note following RCW 69.25.020.

69.25.070 Egg handler’s or dealer’s license—Denial, suspension, revocation, or conditional issuance. The department may deny, suspend, revoke, or issue a license or a conditional license if it determines that an applicant or licensee has committed any of the following acts:

(1) That the applicant or licensee is violating or has violated any of the provisions of this chapter or rules and regulations adopted thereunder.

(2) That the application contains any materially false or misleading statement or involves any misrepresentation by any officer, agent, or employee of the applicant.

(3) That the applicant or licensee has concealed or withheld any facts regarding any violation of this chapter by any officer, agent, or employee of the applicant or licensee. [1975 1st ex.s. c 201 § 8.]

69.25.080 Continuous inspection at processing plants—Exemptions—Condemnation and destruction of adulterated eggs and egg products—Reprocessing—Appeal—Inspections of egg handlers. (1) For the purpose of preventing the entry into or movement in intrastate commerce of any egg product which is capable of use as human food and is misbranded or adulterated, the director shall, whenever processing operations are being conducted, unless under inspection by the United States department of agriculture, cause continuous inspection to be made, in accordance with the regulations promulgated under this chapter, of the processing of egg products, in each plant processing egg products for commerce, unless exempted under RCW 69.25.170. Without restricting the application of the preceding sentence to other kinds of establishments within its provisions, any food manufacturing establishment, institution, or restaurant which uses any eggs that do not meet the requirements of RCW 69.25.170(1)(a) in the preparation of any articles for human food, shall be deemed to be a plant processing egg products, with respect to such operations.

(2) The director, at any time, shall cause such retention, segregation, and reinspection as he or she deems necessary of eggs and egg products capable of use as human food in each official plant.

(3) Eggs and egg products found to be adulterated at official plants shall be condemned, and if no appeal be taken from such determination or condemnation, such articles shall be destroyed for human food purposes under the supervision of an inspector: PROVIDED, That articles which may by...
reprocessing be made not adulterated need not be condemned and destroyed if so reprocessed under the supervision of an inspector and thereafter found to be not adulterated. If an appeal be taken from such determination, the eggs or egg products shall be appropriately marked and segregated pending completion of an appeal inspection, which appeal shall be at the cost of the appellant if the director determines that the appeal is frivolous. If the determination of condemnation is sustained, the eggs or egg products shall be destroyed for human food purposes under the supervision of an inspector.

(4) The director shall cause such other inspections to be made of the business premises, facilities, inventory, operations, and records of egg handlers, and the records and inventory of other persons required to keep records under RCW 69.25.140, as he or she deems appropriate (and in the case of shell egg packers, packing eggs for the ultimate consumer, at least once each calendar quarter) to assure that only eggs fit for human food are used for such purpose, and otherwise to assure compliance by egg handlers and other persons with the requirements of RCW 69.25.140, except that the director shall cause such inspections to be made as he or she deems appropriate to assure compliance with such requirements at food manufacturing establishments, institutions, and restaurants, other than plants processing egg products. Representatives of the director shall be afforded access to all such places of business for purposes of making the inspections provided for in this chapter. [2012 c 117 § 346; 1975 1st ex.s. c 201 § 9.]

69.25.090 Sanitary operation of official plants—Inspection refused if requirements not met. (1) The operator of each official plant shall operate such plant in accordance with such sanitary practices and shall have such premises, facilities, and equipment as are required by regulations promulgated by the director to effectuate the purposes of this chapter, including requirements for segregation and disposition of restricted eggs.

(2) The director shall refuse to render inspection to any plant whose premises, facilities, or equipment, or the operation thereof, fail to meet the requirements of this section. [1975 1st ex.s. c 201 § 10.]

69.25.100 Egg products—Pasteurization—Labeling requirements—False or misleading labels or containers—Director may order use of withheld—Hearing, determination, and appeal. (1) Egg products inspected at any official plant under the authority of this chapter and found to be not adulterated shall be pasteurized before they leave the official plant, except as otherwise permitted by regulations of the director, and shall at the time they leave the official plant, be in distinctly legible form on their shipping containers or immediate containers, or both, when required by regulations of the director, the official inspection legend and official plant number, of the plant where the products were processed, and such other information as the director may require by regulations to describe the products adequately and to assure that they will not have false or misleading labeling.

(2) No labeling or container shall be used for egg products at official plants if it is false or misleading or has not been approved as required by the regulations of the director.

If the director has reason to believe that any labeling or the size or form of any container in use or proposed for use with respect to egg products at any official plant is false or misleading in any particular, he or she may direct that such use be withheld unless the labeling or container is modified in such manner as he or she may prescribe so that it will not be false or misleading. If the person using or proposing to use the labeling or container does not accept the determination of the director, such person may request a hearing, but the use of the labeling or container shall, if the director so directs, be withheld pending hearing and final determination by the director. Any such determination by the director shall be conclusive unless, within thirty days after receipt of notice of such final determination, the person adversely affected thereby appeals to the superior court in the county in which such person has its principal place of business. [2012 c 117 § 347; 1975 1st ex.s. c 201 § 11.]

69.25.103 Eggs or egg products—In-state production—Associated commercial egg layer operation compliance with applicable standards. Any egg handler or dealer involved with the in-state production of eggs or egg products only intended for sale outside of the state of Washington must ensure that the associated commercial egg layer operation is in compliance with the applicable standards as provided in RCW 69.25.065. [2011 c 306 § 4.]

Effective date—2011 c 306: See note following RCW 69.25.020.

69.25.107 Commercial egg layer operations—Requirements. (1) All commercial egg layer operations required under RCW 69.25.065 to meet the American humane association facility system plan, or an equivalent to the plan, must also ensure that all hens in the operation are provided with:
(a) No less than one hundred sixteen and three-tenths square inches of space per hen; and
(b) Access to areas for nesting, scratching, and perching.

(2) The requirements of this section apply for any commercial egg layer operation on the same dates that RCW 69.25.065 requires compliance with the American humane association facility system plan or an equivalent to the plan. [2011 c 306 § 5.]

Effective date—2011 c 306: See note following RCW 69.25.020.

69.25.110 Prohibited acts and practices. (1) No person shall buy, sell, or transport, or offer to buy or sell, or offer or receive for transportation, in any business in intrastate commerce any restricted eggs, capable of use as human food, except as authorized by regulations of the director under such conditions as he or she may prescribe to assure that only eggs fit for human food are used for such purpose.

(2) No egg handler shall possess with intent to use, or use, any restricted eggs in the preparation of human food for intrastate commerce except that such eggs may be so possessed and used when authorized by regulations of the director under such conditions as he or she may prescribe to assure that only eggs fit for human food are used for such purpose.

(3) No person shall process any egg products for intrastate commerce at any plant except in compliance with the requirements of this chapter.
(4) No person shall buy, sell, or transport, or offer to buy or sell, or offer or receive for transportation, in intrastate commerce any egg products required to be inspected under this chapter unless they have been so inspected and are labeled and packaged in accordance with the requirements of RCW 69.25.100.

(5) No operator of any official plant shall allow any egg products to be moved from such plant if they are adulterated or misbranded and capable of use as human food.

(6) No person shall:
   (a) Manufacture, cast, print, lithograph, or otherwise make any device containing any official mark or simulation thereof, or any label bearing any such mark or simulation, or any form of official certificate or simulation thereof, except as authorized by the director;
   (b) Forge or alter any official device, mark, or certificate;
   (c) Without authorization from the director, use any official device, mark, or certificate, or simulation thereof, or detach, deface, or destroy any official device or mark; or use any labeling or container ordered to be withheld from use under RCW 69.25.100 after final judicial affirmance of such order or expiration of the time for appeal if no appeal is taken under said section;
   (d) Contrary to the regulations prescribed by the director, fail to use, or to detach, deface, or destroy any official device, mark, or certificate;
   (e) Knowingly possess, without promptly notifying the director or his or her representative, any official device or any counterfeit, simulated, forged, or improperly altered official certificate or any device or label, or any eggs or egg products bearing any counterfeit, simulated, forged, or improperly altered official mark;
   (f) Knowingly make any false statement in any shipper’s certificate or other nonofficial or official certificate provided for in the regulations prescribed by the director;
   (g) Knowingly represent that any article has been inspected or exempted, under this chapter when in fact it has not been so inspected or exempted; and
   (h) Refuse access, at any reasonable time, to any representative of the director, to any plant or other place of business subject to inspection under any provisions of this chapter.

(7) No person, while an official or employee of the state or local governmental agency, or thereafter, shall use to his or her own advantage, or reveal other than to the authorized representatives of the United States government or the state in their official capacity, or as ordered by a court in a judicial proceeding, any information acquired under the authority of this chapter concerning any matter which the originator or relator of such information claims to be entitled to protection as a trade secret. [2012 c 117 § 348; 1975 1st ex.s. c 201 § 12.]

69.25.120 Director to cooperate with other agencies—May conduct examinations. The director shall, whenever he or she determines that it would effectuate the purposes of this chapter, cooperate with any state, federal, or other governmental agencies in carrying out any provisions of this chapter. In carrying out the provisions of this chapter, the director may conduct such examinations, investigations, and inspections as he or she determines practicable through any officer or employee of any such agency commissioned by him or her for such purpose. [2012 c 117 § 349; 1975 1st ex.s. c 201 § 13.]

69.25.130 Eggs or egg products not intended for use as human food—Identification or denaturing required. Inspection shall not be provided under this chapter at any plant for the processing of any egg products which are not intended for use as human food, but such articles, prior to their offer for sale or transportation in intrastate commerce, shall be denatured or identified as prescribed by regulations of the director to deter their use for human food. No person shall buy, sell, or transport or offer to buy or sell, or offer or receive for transportation, in intrastate commerce, any restricted eggs or egg products which are not intended for use as human food unless they are denatured or identified as required by the regulations of the director. [1975 1st ex.s. c 201 § 14.]

69.25.140 Records required, access to and copying of. For the purpose of enforcing the provisions of this chapter and the regulations promulgated thereunder, all persons engaged in the business of transporting, shipping, or receiving any eggs or egg products in intrastate commerce or in interstate commerce, or holding such articles so received, and all egg handlers, shall maintain such records showing, for such time and in such form and manner, as the director may prescribe, to the extent that they are concerned therewith, the receipt, delivery, sale, movement, and disposition of all eggs and egg products handled by them, and shall, upon the request of the director, permit him or her at reasonable times to have access to and to copy all such records. [2012 c 117 § 350; 1975 1st ex.s. c 201 § 15.]

69.25.150 Penalties—Liability of employer—Defense. (1)(a) Except as provided in (b) of this subsection, any person violating any provision of this chapter or any rule adopted under this chapter is guilty of a misdemeanor.

   (b) A second or subsequent violation is a gross misdemeanor. Any offense committed more than five years after a previous conviction shall be considered a first offense.

   (2) Whenever the director finds that a person has committed a violation of any of the provisions of this chapter, and that violation has not been punished pursuant to subsection (1) of this section, the director may impose upon and collect from the violator a civil penalty not exceeding one thousand dollars per violation per day. Each violation shall be a separate and distinct offense.

   (3) When construing or enforcing the provisions of RCW 69.25.110, the act, omission, or failure of any person acting for or employed by any individual, partnership, corporation, or association within the scope of the person’s employment or office shall in every case be deemed the act, omission, or failure of such individual, partnership, corporation, or association, as well as of such person.

   (4) No carrier or warehouse operator shall be subject to the penalties of this chapter, other than the penalties for violation of RCW 69.25.140, or 69.25.155, by reason of his or her receipt, carriage, holding, or delivery, in the usual course of business, as a carrier or warehouse operator of eggs or egg products owned by another person unless the carrier or ware-
69.25.155 Interference with person performing official duties.

(1) Notwithstanding any other provision of law, any person who forcibly assaults, resists, impedes, intimidates, or interferes with any person while engaged in or on account of the performance of his or her official duties under this chapter is guilty of a class C felony and shall be punished by a fine of not more than five thousand dollars or imprisonment in a state correctional facility for not more than three years, or both.

(2) Whoever, in the commission of any act described in subsection (1) of this section, uses a deadly or dangerous weapon is guilty of a class B felony and shall be punished by a fine of not more than ten thousand dollars or by imprisonment in a state correctional facility for not more than ten years, or both. [2003 c 53 § 318.]

69.25.160 Notice of violation—May take place of prosecution. Before any violation of this chapter, other than RCW 69.25.155, is reported by the director to any prosecuting attorney for institution of a criminal proceeding, the person against whom such proceeding is contemplated shall be given reasonable notice of the alleged violation and opportunity to present his or her views orally or in writing with regard to such contemplated proceeding. Nothing in this chapter shall be construed as requiring the director to report for criminal prosecution violation of this chapter whenever he or she believes that the public interest will be adequately served and compliance with this chapter obtained by a suitable written notice of warning. [2003 c 53 § 319; 1975 1st ex.s. c 201 § 17.]

69.25.170 Exemptions permitted by rule of director.

(1) The director may, by regulation and under such conditions and procedures as he or she may prescribe, exempt from specific provisions of this chapter:

(a) The sale, transportation, possession, or use of eggs which contain no more restricted eggs than are allowed by the tolerance in the official state standards for consumer grades for shell eggs;

(b) The processing of egg products at any plant where the facilities and operating procedures meet such sanitary standards as may be prescribed by the director, and where the eggs received or used in the manufacture of egg products contain no more restricted eggs than are allowed by the official standards of the state consumer grades for shell eggs, and the egg products processed at such plant;

(c) The sale of eggs by any poultry producer from his or her own flocks directly to a household consumer exclusively for use by such consumer and members of his or her household and his or her nonpaying guests and employees, and the transportation, possession, and use of such eggs in accordance with this subsection;

(d) The sale of eggs by shell egg packers on his or her own premises directly to household consumers for use by such consumer and members of his or her household and his or her nonpaying guests and employees, and the transportation, possession, and use of such eggs in accordance with this subsection;

(e) The sale of eggs by any egg producer with an annual egg production from a flock of three thousand hens or less.

(2) The director may modify or revoke any regulation granting exemption under this chapter whenever he or she deems such action appropriate to effectuate the purposes of this chapter. [2012 c 117 § 351; 1995 c 374 § 28; 1975 1st ex.s. c 201 § 18.]

Additional notes found at www.leg.wa.gov

69.25.180 Limiting entry of eggs and egg products into official plants. The director may limit the entry of eggs and egg products and other materials into official plants under such conditions as he or she may prescribe to assure that allowing the entry of such articles into such plants will be consistent with the purposes of this chapter. [2012 c 117 § 352; 1975 1st ex.s. c 201 § 19.]

69.25.190 Embargo of eggs or egg products in violation of this chapter—Time limit—Removal of official marks. Whenever any eggs or egg products subject to this chapter are found by any authorized representative of the director upon any premises and there is reason to believe that they are or have been processed, bought, sold, possessed, used, transported, or offered or received for sale or transportation in violation of this chapter, or that they are in any other way in violation of this chapter, or whenever any restricted eggs capable of use as human food are found by such a representative in the possession of any person not authorized to acquire such eggs under the regulations of the director, such articles may be embargoed by such representative for a reasonable period but not to exceed twenty days, pending action under RCW 69.25.200 or notification of any federal or other governmental authorities having jurisdiction over such articles, and shall not be moved by any person from the place at which they are located when so detained until released by such representative. All official marks may be required by such representative to be removed from such articles before they are released unless it appears to the satisfaction of the director that the articles are eligible to retain such marks. [1975 1st ex.s. c 201 § 20.]

69.25.200 Embargo—Petition for court order affirming—Removal of embargo or destruction or correction and release—Court costs, fees, administrative expenses—
Bond may be required. When the director has embargoed any eggs or egg products, he or she shall petition the superior court of the county in which the eggs or egg products are located for an order affirming such embargo. Such court shall have jurisdiction for cause shown and after a prompt hearing to any claimant of eggs or egg products, shall issue an order which directs the removal of such embargo or the destruction or correction and release of such eggs and egg products. An order for destruction or the correction and release of such eggs and egg products shall contain such provisions for the payment of pertinent court costs and fees and administrative expenses as is equitable and which the court deems appropriate in the circumstances. An order for correction and release may contain such provisions for a bond as the court finds indicated in the circumstance. [2012 c 117 § 353; 1975 1st ex.s. c 201 § 21.]

69.25.210 Embargo—Order affirming not required, when. The director need not petition the superior court as provided for in RCW 69.25.200 if the owner or claimant of such eggs or egg products agrees in writing to the disposition of such eggs or egg products as the director may order. [1975 1st ex.s. c 201 § 22.]

69.25.220 Embargo—Consolidation of petitions. Two or more petitions under RCW 69.25.200 which pend at the same time and which present the same issue and claimant hereunder may be consolidated for simultaneous determination by one court of competent jurisdiction, upon application to any court of jurisdiction by the director or claimant. [1975 1st ex.s. c 201 § 23.]

69.25.230 Embargo—Sampling of article. The claimant in any proceeding by petition under RCW 69.25.200 shall be entitled to receive a representative sample of the article subject to such proceedings upon application to the court of competent jurisdiction made at any time after such petition and prior to the hearing thereon. [1975 1st ex.s. c 201 § 24.]

69.25.240 Condemnation—Recovery of damages restricted. No state court shall allow the recovery of damages for administrative action for condemnation under the provisions of this chapter, if the court finds that there was probable cause for such action. [1975 1st ex.s. c 201 § 25.]

69.25.250 Assessment—Rate, applicability, time of payment—Reports—Contents, frequency—Exemption. (1)(a) There is hereby levied an assessment not to exceed three mills per dozen eggs entering intrastate commerce, as prescribed by rules issued by the director. The assessment is applicable to all eggs entering intrastate commerce, except as provided in RCW 69.25.170 and 69.25.290, and must be paid to the director on a monthly basis on or before the tenth day following the month the eggs enter intrastate commerce.

(b) The director may require reports by egg handlers or dealers along with the payment of the assessment fee. The reports may include any and all pertinent information necessary to carry out the purposes of this chapter.

(c) The director may, by rule, require egg container manufacturers to report on a monthly basis all egg containers sold to any egg handler or dealer and bearing such egg handler or dealer’s permanent number.

(2) Egg products in intrastate commerce are exempt from the assessment in subsection (1) of this section. [2011 c 306 § 6; 1995 c 374 § 29; 1993 sp.s. c 19 § 12; 1975 1st ex.s. c 201 § 26.]

Effective date—2011 c 306: See note following RCW 69.25.020.

Additional notes found at www.leg.wa.gov

69.25.260 Assessment—Prepayment by purchase of egg seals—Permit for printing seal on containers or labels. Any egg handler or dealer may prepay the assessment provided for in RCW 69.25.250 by purchasing Washington state egg seals from the director to be placed on egg containers showing that the proper assessment has been paid. Any carton manufacturer or printer may apply to the director for a permit to place reasonable facsimiles of the Washington state egg seals to be imprinted on egg containers or on the identification labels which show egg grade and size and the name of the egg handler or dealer. The director shall, from time to time, prescribe rules and regulations governing the affixing of seals and he or she is authorized to cancel any such permit issued pursuant to this chapter, whenever he or she finds that a violation of the terms under which the permit has been granted has been violated. [2012 c 117 § 354; 1979 ex.s. c 238 § 10; 1975 1st ex.s. c 201 § 27.]

Additional notes found at www.leg.wa.gov

69.25.270 Assessment—Monthly payment—Audit—Failure to pay, penalty. Every egg handler or dealer who pays assessments required under the provisions of this chapter on a monthly basis in lieu of seals shall be subject to audit by the director at such frequency as is deemed necessary by the director. The cost to the director for performing such audit shall be chargeable to and payable by the egg handler or dealer subject to audit. Failure to pay assessments when due or refusal to pay for audit costs may be cause for a summary suspension of an egg handler’s or dealer’s license and a charge of one percent per month, or fraction thereof shall be added to the sum due the director, for each remittance not received by the director when due. The conditions and charges applicable to egg handlers and dealers set forth herein shall also be applicable to payments due the director for facsimiles of seals placed on egg containers. [1987 c 393 § 16; 1975 1st ex.s. c 201 § 28.]

69.25.280 Assessment—Use of proceeds. The proceeds from assessment fees paid to the director shall be retained for the inspection of eggs and carrying out the provisions of this chapter relating to eggs. [1975 1st ex.s. c 201 § 29.]

69.25.290 Assessment—Exclusions. The assessments provided in this chapter shall not apply to:

(1) Sale and shipment to points outside of this state;

(2) Sale to the United States government and its instrumentalities;

(3) Sale to breaking plants for processing into egg products;

(4) Sale between egg dealers. [1975 1st ex.s. c 201 § 30.]
69.25.300 Transfer of moneys in state egg account. All moneys in the state egg account, created by *RCW 69.24.450, at the time of July 1, 1975, shall be transferred to the director and shall be retained and expended for administering and carrying out the purposes of this chapter. [1975 1st ex.s. c 201 § 31.]

*Reviser’s note:* RCW 69.24.450 was repealed by 1975 1st ex.s. c 201 § 40.

69.25.310 Containers—Marking required—Obliteration of previous markings required for reuse—Temporary use of another handler’s or dealer’s permanent number—Penalty. (1) All containers used by an egg handler or dealer to package eggs shall bear the name and address or the permanent number issued by the director to said egg handler or dealer. Such permanent number shall be displayed in a size and location prescribed by the director. It shall be a violation for any egg handler or dealer to use a container that bears the permanent number of another egg handler or dealer unless such number is totally obliterated prior to use. The director may in addition require the obliteration of any or all markings that may be on any container which will be used for eggs by an egg handler or dealer.

(2) Notwithstanding subsection (1) of this section and following written notice to the director, licensed egg handlers and dealers may use new containers bearing an another handler’s or dealer’s permanent number on a temporary basis, in any event not longer than one year, with the consent of such other handler or dealer for the purpose of using up existing container stocks. Sale of container stock shall constitute agreement by the parties to use the permanent number. [1995 c 374 § 30; 1975 1st ex.s. c 201 § 32.]

Additional notes found at www.leg.wa.gov

69.25.320 Records required, additional—Sales to retailer or food service—Exception—Defense to charged violation—Sale of eggs deteriorated due to storage time—Requirements for storage, display, or transportation. (1) In addition to any other records required to be kept and furnished the director under the provisions of this chapter, the director may require any person who sells to any retailer, or to any restaurant, hotel, boarding house, bakery, or any institution or concern which purchases eggs for serving to guests or patrons thereof or for its use in preparation of any food product for human consumption, candled or graded eggs other than those of his or her own production sold and delivered on the premises where produced, to furnish that retailer or other purchaser with an invoice covering each such sale, showing the exact grade or quality, and the size or weight of the eggs sold, according to the standards prescribed by the director, together with the name and address of the person by whom the eggs were sold. The person selling and the retailer or other purchaser shall keep a copy of said invoice on file at his or her place of business for a period of thirty days, during which time the copy shall be available for inspection at all reasonable times by the director: PROVIDED, That no retailer or other purchaser shall be guilty of a violation of this chapter if he or she can establish a guarantee from the person from whom the eggs were purchased to the effect that they, at the time of purchase, conformed to the information required by the director on such invoice: PROVIDED FURTHER, that if the retailer or other purchaser having labeled any such eggs in accordance with the invoice keeps them for such a time after they are purchased as to cause them to deteriorate to a lower grade or standard, and sells them under the label of the invoice grade or standard, he or she shall be guilty of a violation of this chapter.

(2) Each retailer and each distributor shall store shell eggs awaiting sale or display eggs under clean and sanitary conditions in areas free from rodents and insects. Shell eggs must be stored up off the floor away from strong odors, pesticides, and cleaners.

(3) After being received at the point of first purchase, all graded shell eggs packed in containers for the purpose of sale to consumers shall be held and transported under refrigeration at ambient temperatures no greater than forty-five degrees Fahrenheit (seven and two-tenths degrees Celsius). This provision shall apply without limitation to retailers, institutional users, dealer/wholesalers, food handlers, transportation firms, or any person who handles eggs after the point of first purchase.

(4) No invoice shall be required on eggs when packed for sale to the United States department of defense, or a component thereof, if labeled with grades promulgated by the United States secretary of agriculture. [2012 c 117 § 355; 1995 c 374 § 31; 1975 1st ex.s. c 201 § 33.]

Additional notes found at www.leg.wa.gov

69.25.900 Savings. The enactment of this chapter shall not have the effect of terminating or in any way modifying any liability, civil or criminal, which shall already be in existence on July 1, 1975. [1975 1st ex.s. c 201 § 35.]

69.25.910 Chapter is cumulative and nonexclusive. The provisions of this chapter shall be cumulative and nonexclusive and shall not affect any other remedy at law. [1975 1st ex.s. c 201 § 37.]

69.25.920 Severability—1975 1st ex.s. c 201. If any provision of this chapter, or its application to any person or circumstance is held invalid, the remainder of the chapter, or the application of the provision to other persons or circumstances is not affected. [1975 1st ex.s. c 201 § 38.]

69.25.930 Short title. This act may be known and cited as the "Washington wholesome eggs and egg products act". [1975 1st ex.s. c 201 § 39.]

Chapter 69.28 RCW

HONEY

Sections
69.28.020 Enforcement power and duty of director and agents.
69.28.025 Rules and regulations have force of law.
69.28.030 Rules prescribing standards.
69.28.040 Right to enter, inspect, and take samples.
69.28.050 Containers to be labeled.
69.28.060 Requisites of markings.
69.28.070 "Marked" defined—When honey need not be marked.
69.28.080 Purchaser to be advised of standards—Exceptions.
69.28.090 Forgery, simulation, etc., unlawful.
69.28.095 Unlawful mutilation or removal of seals, marks, etc., used by director.
69.28.100 Marks for "slack-filled" container.
69.28.110 Use of used containers.

[Title 69 RCW—page 38] 2012
69.28.020 Enforcement power and duty of director and agents. The director is hereby empowered, through his or her duly authorized agents, to enforce all provisions of this chapter. The director shall have the power to define, promulgate, and enforce such reasonable regulations as he or she may deem necessary in carrying out the provisions of this chapter. [2012 c 117 § 356; 1939 c 199 § 29; RRS § 6163-29. FORMER PART OF SECTION: 1939 c 199 § 44 now codified as RCW 69.28.025.]

69.28.025 Rules and regulations have force of law. Any rules or regulations promulgated and published by the director under the provisions of this chapter shall have the force and effect of law. [1939 c 199 § 44; RRS § 6163-44. Formerly RCW 69.28.020, part.]

69.28.030 Rules prescribing standards. The director is hereby authorized, and it shall be his or her duty, upon the taking effect of this chapter and from time to time thereafter, to adopt, establish, and promulgate reasonable rules and regulations specifying grades or standards of quality governing the sale of honey: PROVIDED, That, in the interest of uniformity, such grades and standards of quality shall conform as nearly to those established by the United States department of agriculture as local conditions will permit. [2012 c 117 § 357; 1939 c 199 § 24; RRS § 6163-24.]

69.28.040 Right to enter, inspect, and take samples. The director or any of his or her duly authorized agents shall have the power to enter and inspect at reasonable times every place, vehicle, plant, or other place where honey is being produced, stored, packed, transported, exposed, or offered for sale, and to inspect all such honey and the containers thereof and to take for inspection such samples of said honey as may be necessary. [2012 c 117 § 358; 1939 c 199 § 28; RRS § 6163-28.]

69.28.050 Containers to be labeled. It shall be unlawful to deliver for shipment, ship, transport, sell, expose or offer for sale any containers or subcontainers of honey within this state unless they shall be conspicuously marked with the name and address of the producer or distributor, the net weight of the honey, the grade of the honey, and, if imported from any foreign country, the name of the country or territory from which the said honey was imported, or if a blend of honey, any part of which is foreign honey, the container must be labeled with the name of the country or territory where such honey was produced and the proportion of each foreign honey used in the blend. [1939 c 199 § 32; RRS § 6163-32.]

69.28.060 Requisites of markings. When any markings are used or required to be used under this chapter on any container of honey to identify the container or describe the contents thereof, such markings must be plainly and conspicuously marked, stamped, stenciled, printed, labeled or branded in the English language, in letters large enough to be discernible by any person, on the front, side or top of any container. [1939 c 199 § 35; RRS § 6163-35.]

69.28.070 "Marked" defined—When honey need not be marked. The term "marked" shall mean printed in the English language on the top, front or side of any container containing honey: PROVIDED, That it shall not be necessary to mark honey sold by the producer thereof to any distributor, packer or manufacturer with the net weight, color or grade if the honey is to be used in the manufacture of honey products or is to be graded and packaged by the distributor or packer for resale. [1939 c 199 § 21; RRS § 6163-21.]

69.28.080 Purchaser to be advised of standards—Exceptions. It shall be unlawful for any person to deliver, sell, offer, or expose for sale any honey for human consumption within the state without notifying the person or persons purchasing or intending to purchase the same, of the exact grade or quality of such honey, according to the standards prescribed by the director, by stamping or printing on the container of any such honey such grade or quality: PROVIDED, This section shall not apply to honey while it is in transit in intrastate commerce from one establishment to the other, to be processed, labeled, or repacked. [1961 c 60 § 1; 1957 c 103 § 1; 1949 c 105 § 6; 1939 c 199 § 39; Rem. Supp. 1949 § 6163-39.]

69.28.090 Forgery, simulation, etc., of marks, labels, etc., unlawful. It shall be unlawful to forge, counterfeit, simulate, falsely represent or alter without proper authority any mark, stamp, tab, label, seal, sticker or other identification device provided by this chapter. [1961 c 60 § 2; 1939 c 199 § 40; RRS § 6163-40. FORMER PART OF SECTION: 1939 c 199 § 41 now codified as RCW 69.28.095.]
69.28.095 Unlawful mutilation or removal of seals, marks, etc., used by director. It shall be unlawful to mutilate, destroy, obliterate, or remove without proper authority, any mark, stamp, tag, label, seal, sticker or other identification device used by the director under the provisions of this chapter. [1939 c 199 § 41; RRS § 6163-41. Formerly RCW 69.28.090, part.]

69.28.100 Marks for "slack-filled" container. Any slack-filled container shall be conspicuously marked "slack-filled". [1939 c 199 § 36; RRS § 6163-36. FORMER PART OF SECTION: 1939 c 199 § 10 now codified as RCW 69.28.270.]

69.28.110 Use of used containers. It shall be unlawful to sell, offer, or expose for sale to the consumer any honey in any secondhand or used containers which formerly contained honey, unless all markings as to grade, name and weight have been obliterated, removed or erased. [1939 c 199 § 37; RRS § 6163-37.]

69.28.120 Floral source labels. Any honey which is a blend of two or more floral types of honey shall not be labeled as a honey product from any one particular floral source alone. [1939 c 199 § 34; RRS § 6163-34.]

69.28.130 Adulterated honey—Sale or offer unlawful. It shall be unlawful for any person to sell, offer or intend for sale any adulterated honey as honey. [1939 c 199 § 26; RRS § 6163-26. FORMER PART OF SECTION: 1939 c 199 §§ 27 and 33 now codified as RCW 69.28.133 and 69.28.135.]

69.28.133 Nonconforming honey—Sale or offer unlawful. It shall be unlawful for any person to sell, offer or intend for sale any honey which does not conform to the provisions of this chapter or any regulation promulgated by the director under this chapter. [1939 c 199 § 27; RRS § 6163-27. Formerly RCW 69.28.130, part.]

69.28.135 Warning-tagged honey—Movement prohibited. It shall be unlawful to move any honey or containers of honey to which any warning tag or notice has been affixed except under authority from the director. [1939 c 199 § 33; RRS § 6163-33. Formerly RCW 69.28.130, part.]

69.28.140 Possession of unlawful honey as evidence. Possession by any person, of any honey which is sold, exposed or offered for sale in violation of this chapter shall be prima facie evidence that the same is kept or shipped to the said person, in violation of the provisions of this chapter. [1939 c 199 § 30; RRS § 6163-30.]

69.28.170 Inspectors—Prosecutions. It shall be the duty of the director to enforce this chapter and to appoint and employment [employ] such inspectors as may be necessary therefor. The director shall notify the prosecuting attorneys for the counties of the state of violations of this chapter occurring in their respective counties, and it shall be the duty of the respective prosecuting attorneys immediately to institute and prosecute proceeding in their respective counties and to enforce the penalties provided for by this chapter. [1939 c 199 § 43; RRS § 6163-43.]

69.28.180 Violation of rules and regulations unlawful. It shall be unlawful for any person to violate any rule or regulation promulgated by the director under the provisions of this chapter. [1939 c 199 § 25; RRS § 6163-25. FORMER PART OF SECTION: 1939 c 199 § 44 now codified in RCW 69.28.185.]

69.28.185 Penalty. Any person who violates any of the provisions of this chapter shall be guilty of a misdemeanor, and upon violation thereof shall be punishable by a fine of not more than five hundred dollars or imprisonment in the county jail for a period of not more than six months or by both such fine and imprisonment. [1939 c 199 § 42; RRS § 6163-42. Formerly RCW 69.28.180, part.]

69.28.190 "Director" defined. The term "director" means the director of agriculture of the state of Washington or his or her duly authorized representative. [2012 c 117 § 359; 1939 c 199 § 2; RRS § 6163-2. Formerly RCW 69.28.010, part.]

69.28.200 "Container" defined. The term "container" shall mean any box, crate, chest, carton, barrel, keg, bottle, jar, can or any other receptacle containing honey. [1939 c 199 § 3; RRS § 6163-3.]

69.28.210 "Subcontainer" defined. The term "sub-container" shall mean any section box or other receptacle used within a container. [1939 c 199 § 4; RRS § 6163-4.]

69.28.220 "Section box" defined. The term "section box" shall mean the wood or other frame in which bees have built a small comb of honey. [1939 c 199 § 5; RRS § 6163-5.]

69.28.230 "Clean and sound containers" defined. The term "clean and sound containers" shall mean containers which are virtually free from rust, stains or leaks. [1939 c 199 § 6; RRS § 6163-6.]

69.28.240 "Pack," "packing," or "packed" defined. The term "pack", "packing", or "packed" shall mean the arrangement of all or part of the subcontainers in any container. [1939 c 199 § 7; RRS § 6163-7.]

69.28.250 "Label" defined. The term "label" shall mean a display of written, printed or graphic matter upon the immediate container of any article. [1939 c 199 § 8; RRS § 6163-8.]

69.28.260 "Person" defined. The term "person" includes individual, partnership, corporation and/or association. [1939 c 199 § 9; RRS § 6163-9.]

69.28.270 "Slack-filled" defined. The term "slack-filled" shall mean that the contents of any container occupy less than ninety-five percent of the volume of the closed con-
condition, or in any other respects, to any of that which is surface, honey which is so superior in quality, appearance or condition, or in any other respects, to any of that which is concealed or unexposed as to materially misrepresent any part of the lot, load, arrangement or display.  

Deceptive arrangement defined. The term "deceptive arrangement" shall mean any lot or load, part of the lot, load, arrangement or display.  

Mislabeled defined. The term "mislabeled" shall mean the placing or presence of any false or misleading statement, design or device upon, or in connection with, any container or lot of honey, or upon the label, lining or wrapper of any such container, or any placard used in connection therewith, and having reference to such honey. A statement, design or device is false and misleading when the honey to which it refers does not conform in every respect to such statement.  

Placard defined. The term "placard" means any sign, label or designation, other than an oral designation, used with any honey as a description or identification thereof.  

Honey defined. The term "honey" as used herein is the nectar of floral exudations of plants, gathered and stored in the comb by honey bees (apis mellifica). It is laevorotatory, contains not more than twenty-five percent of water, not more than twenty-five one-hundredths of one percent of ash, not more than eight percent of sucrose, its specific gravity is 1.412, its weight not less than eleven pounds twelve ounces per standard gallon of 231 cubic inches at sixty-eight degrees Fahrenheit.  

Comb-honey defined. The term "comb-honey" means honey which has not been extracted from the comb.  

Extracted honey defined. The term "extracted honey" means honey which has been removed from the comb.  

Crystallized honey defined. The term "crystallized honey" means honey which has assumed a solid form due to the crystallization of one or more of the natural sugars therein.  

Honeydew defined. The term "honeydew" is the saccharine exudation of plants, other than nectarous exudations, gathered and stored in the comb by honey bees (apis mellifica) and is dextrorotatory.  

Foreign material defined. The term "foreign material" means pollen, wax particles, insects, or materials not deposited by bees.  

Foreign honey defined. The term "foreign honey" means any honey not produced within the continental United States.  

Adulterated honey defined. The term "adulterated honey" means any honey to which has been added honeydew, glucose, dextrose, molasses, sugar, sugar syrup, invert sugar, or any other similar product or products, other than the nectar of floral exudations of plants gathered and stored in the comb by honey bees.  

Serious damage defined. The term "serious damage" means any injury or defect that seriously affects the edibility or shipping quality of the honey.  

Labeling requirements for artificial honey or mixtures containing honey. (1) No person shall sell, keep for sale, expose or offer for sale, any article or product in imitation or semblance of honey branded exclusively as "honey", "liquid or extracted honey", "strained honey" or "pure honey".  

(2) No person, firm, association, company or corporation shall manufacture, sell, expose or offer for sale, any compound or mixture branded or labeled exclusively as honey which shall be made up of honey mixed with any other substance or ingredient.  

(3) Whenever honey is mixed with any other substance or ingredient and the commodity is to be marketed in imitation or semblance of honey, the product shall be labeled with the word "artificial" or "imitation" in the same type size and style as the word "honey";  

(4) Whenever any substance or commodity is to be marketed in imitation or semblance of honey, but contains no honey, the product shall not be branded or labeled with the word "honey" and/or depict thereon a picture or drawing of a bee, bee hive, or honeycomb;  

(5) Whenever honey is mixed with any other substance or ingredient and the commodity is to be marketed, there shall be printed on the package containing such compound or mixture a statement giving the ingredients of which it is made; if honey is one of such ingredients it shall be so stated in the same size type as are the other ingredients; nor shall such compound or mixture be branded or labeled exclusively with the word "honey" in any form other than as herein provided; nor shall any product in semblance of honey, whether a mixture or not, be sold, exposed or offered for sale as honey, or branded or labeled exclusively with the word "honey", unless such article is pure honey.  

Embargo on honey or product—Notice by director—Removal. Whenever the director shall find, or shall have probable cause to believe, that any honey or product subject to the provisions of this chapter, as now or hereafter amended, is in intrastate commerce, which was introduced into such intrastate commerce in violation of the provisions of this chapter, as now or hereafter amended, he or she is hereby authorized to affix to such honey or product a notice placing an embargo on such honey or product, and prohib-
ing its sale in intrastate commerce, and no person shall move or sell such honey or product without first receiving permission from the director to move or sell such honey or product. But if, after such honey or product has been embargoed, the director shall find that such honey or product does not involve a violation of this chapter, as now or hereafter amended, such embargo shall be forthwith removed. [2012 c 117 § 360; 1975 1st ex.s. c 283 § 3.]

69.28.420 Embargo on honey or product—Court order affirming, required—Order for destruction or correction and release—Bond. When the director has embargoed any honey or product, he or she shall, no later than twenty days after the affixing of notice of its embargo, petition the superior court for an order affirming such embargo. Such court shall then have jurisdiction, for cause shown and after prompt hearing to any claimant of such honey or product, to issue an order which directs the removal of such embargo or the destruction or the correction and release of such honey or product. An order for destruction or correction and release shall contain such provision for the payment of pertinent court costs and fees and administrative expenses, as is equitable and which the court deems appropriate in the circumstances. An order for correction and release may contain such provision for bond, as the court finds indicated in the circumstances. [2012 c 117 § 361; 1975 1st ex.s. c 283 § 4.]

69.28.430 Consolidation of petitions presenting same issue and claimant. Two or more petitions under this chapter, as now or hereafter amended, which pend at the same time and which present the same issue and claimant hereunder, shall be consolidated for simultaneous determination by one court of jurisdiction, upon application to any court of jurisdiction by the director or by such claimant. [1975 1st ex.s. c 283 § 5.]

69.28.440 Sample of honey or product may be obtained—Procedure. The claimant in any proceeding by petition under this chapter, as now or hereafter amended, shall be entitled to receive a representative sample of the honey or product subject to such proceeding, upon application to the court of jurisdiction made at any time after such petition and prior to the hearing thereon. [1975 1st ex.s. c 283 § 6.]

69.28.450 Recovery of damages barred if probable cause for embargo. No state court shall allow the recovery of damages for embargo under this chapter, as now or hereafter amended, if the court finds that there was probable cause for such action. [1975 1st ex.s. c 283 § 7.]

69.28.900 Severability—1939 c 199. If any provisions of this chapter, or the application thereof to any person or circumstance, is held invalid, the remainder of the chapter, and the application of such provisions to other persons or circumstances, shall not be affected thereby. If any section, subsection, sentence, clause, or phrase of this chapter is for any reason held to be unconstitutional, such decisions shall not affect the validity of the remaining portions of this chapter. The legislature hereby declares that it would have passed this chapter and each section, subsection, sentence, clause and phrase thereof, irrespective of the fact that any one or more of the other sections, subsections, sentences, clauses and phrases be declared unconstitutional. [1939 c 199 § 45; RRS § 6163-45.]

69.28.910 Short title. This chapter may be known and cited as the Washington state honey act. [1939 c 199 § 1; RRS § 6163-1.]

Chapter 69.30 RCW
SANITARY CONTROL OF SHELLFISH

Sections
69.30.005 Purpose.
69.30.010 Definitions.
69.30.020 Approved shellfish tag or label—Requirement to sell or offer to sell shellfish.
69.30.030 Rules and regulations—Duties of state board of health.
69.30.050 Shellfish growing areas—Requirements to harvest—Certificates of approval.
69.30.060 Certificates of approval—Culling, shucking, packing establishments.
69.30.070 Certificates of approval—Compliance with other laws and rules required.
69.30.080 Licenses or certificates of approval—Department may deny, revoke, or suspend.
69.30.085 License, certificate of approval—Denial, revocation, suspension—Prohibited acts—Penalties.
69.30.110 Possession or sale in violation of chapter—Enforcement—Seizure—Disposal.
69.30.120 Inspection by department—Access to regulated business or entity—Administrative inspection warrant.
69.30.130 Water pollution laws and rules applicable.
69.30.140 Penalties.
69.30.145 Civil penalties.
69.30.150 Civil penalties—General provisions.
69.30.900 Severability—1955 c 144.

Shellfish: Chapter 77.60 RCW.

69.30.005 Purpose. The purpose of this chapter is to provide for the sanitary control of shellfish. Protection of the public health requires assurances that commercial shellfish are harvested only from approved growing areas and that processing of shellfish is conducted in a safe and sanitary manner. [1989 c 200 § 2.]

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

(1) "Approved shellfish tag or label" means a tag or label meeting the requirements of the national shellfish sanitation program model ordinance.

(2) "Commercial quantity" means any quantity exceeding: (a) Forty pounds of mussels; (b) one hundred oysters; (c) fourteen horse clams; (d) six geoducks; (e) fifty pounds of hard or soft shell clams; or (f) fifty pounds of scallops. The poundage in this subsection (2) constitutes weight with the shell.

(3) "Department" means the state department of health.

(4) "Establishment" means the buildings, together with the necessary equipment and appurtenances, used for the storage, culling, shucking, packing and/or shipping of shellfish in commercial quantity or for sale for human consumption.

(5) "Ex officio fish and wildlife officer" means an ex officio fish and wildlife officer as defined in RCW 77.08.010.
69.30.020 Approved shellfish tag or label—Requirement to sell or offer to sell shellfish. It is unlawful to sell or offer to sell shellfish in this state unless the shellfish bear an approved shellfish tag or label indicating compliance with the sanitary requirements of this state or a state, territory, province, or country of origin whose requirements are equal or comparable to those established pursuant to this chapter. The department, a fish and wildlife officer, or an ex officio fish and wildlife officer may immediately seize containers of shellfish that are not affixed with an approved shellfish tag or label. [2011 c 194 § 2; 1955 c 144 § 2.]

69.30.030 Rules and regulations—Duties of state board of health. (1) The state board of health shall adopt rules governing the sanitation of shellfish, shellfish growing areas, and shellfish plant facilities and operations in order to protect public health and carry out the provisions of this chapter. Such rules and regulations may include reasonable sanitary requirements relative to the quality of shellfish growing waters and areas, boat and barge sanitation, building construction, water supply, sewage and waste water disposal, lighting and ventilation, insect and rodent control, shell disposal, garbage and waste disposal, cleanliness of establishment, the handling, storage, construction and maintenance of equipment, the handling, storage and refrigeration of shellfish, the identification of containers, and the handling, maintenance, and storage of permits, certificates, and records regarding shellfish taken under this chapter. The state board of health shall adopt rules governing procedures for the disposition of seized shellfish.

(2) The state board of health shall consider the most recent version of the national shellfish sanitation program model ordinance, adopted by the interstate shellfish sanitation conference, when adopting rules. [2011 c 194 § 3; 1995 c 147 § 2; 1955 c 144 § 3.]

69.30.050 Shellfish growing areas—Requirements to harvest—Certificates of approval. (1) It is unlawful for a person to harvest shellfish from shellfish growing areas in a commercial quantity or for sale for human consumption unless the shellfish growing area:

(a) Has a valid certificate of approval; and

(b) Meets the requirements of this chapter and the rules adopted under this chapter.

(2) A person may not remove shellfish in a commercial quantity or for sale for human consumption from a shellfish growing area in the state of Washington unless:

(a) The person has received a certificate of approval for the shellfish growing area from the department; and

(b) Approved shellfish tags are affixed to each container of shellstock prior to removal from the shellfish growing area, except bulk tagging is permitted as allowed in the national shellfish sanitation program model ordinance.

(3) Before issuing a certificate of approval, the department shall inspect the shellfish growing area. The department shall issue a certificate of approval if the area meets the requirements of this chapter and the rules adopted under this chapter.

(4) A certificate of approval is valid for a period of twelve months. The department may revoke a certificate of approval at any time the area is found out of compliance with the requirements of this chapter or the rules adopted under this chapter.

(5) It is unlawful to remove shellfish from shellfish growing areas without a certificate of approval in a commercial quantity for purposes other than human consumption, including but not limited to use as bait or seed, unless:

(a) The shellfish operation and shellfish growing area is readily available to monitoring and inspections; and

(b) The department has determined the shellfish operation is designed to ensure that shellfish harvested from such an area is not diverted for human consumption.

(6) Nothing in this section prohibits a person from removing shellfish for use as bait or seed from an approved shellfish growing area.

(7) The department’s certificate of approval to harvest shellfish for purposes other than human consumption shall specify:

(a) The date or dates and time of harvest;

(b) All applicable conditions of harvest;

(c) Identification by tagging, dying, or other department-approved means; and

(d) Information about the removal method, transportation method, processing technique, sale details, and other factors to ensure that shellfish harvested from such areas are not diverted for human consumption. [2011 c 194 § 4; 1995 c 147 § 3; 1985 c 51 § 2; 1955 c 144 § 5.]

69.30.060 Certificates of approval—Culling, shucking, packing establishments. (1) It is unlawful for a person to cull, shuck, or pack shellfish in the state of Washington in a commercial quantity or for sale for human consumption unless the establishment in which such operations are conducted has been certified by the department as meeting the requirements of the state board of health.

(2) A person may not cull, shuck, or pack shellfish within the state of Washington in a commercial quantity or
for sale for human consumption, unless the person has received a certificate of approval from the department for the establishment in which such operations will be done.

(3) Before issuing a certificate of approval, the department shall inspect the establishment, and if the establishment meets the rules of the state board of health, the department shall issue a certificate of approval. Such certificates of approval shall be issued for a period not to exceed twelve months, and may be revoked at any time the establishment or the operations are found not to be in compliance with the rules of the state board of health. [2011 c 194 § 5; 1985 c 51 § 3; 1955 c 144 § 6.]

69.30.070 Certificates of approval—Compliance with other laws and rules required. Any certificate of approval issued under the provisions of this chapter shall not relieve any person from complying with the laws, rules and/or regulations of the department of fish and wildlife, relative to shellfish. [1994 c 264 § 40; 1955 c 144 § 7.]

69.30.080 Licenses or certificates of approval—Department may deny, revoke, or suspend. (1) The department may deny, revoke, or suspend a person’s license or certificate of approval for:

(a) Violations of this chapter or the rules adopted under this chapter; or
(b) Harassing or threatening an authorized representative of the department during the performance of his or her duties.

(2) RCW 43.70.115 governs notice of a license denial, revocation, suspension, or modification and provides the right to an adjudicative proceeding. [2011 c 194 § 6; 1991 c 3 § 304; 1989 c 175 § 125; 1979 c 141 § 71; 1955 c 144 § 8.]

Additional notes found at www.leg.wa.gov

69.30.085 License, certificate of approval—Denial, revocation, suspension—Prohibited acts—Penalties. (1) A person, or its director or officer, whose license or certificate of approval is denied, revoked, or suspended as a result of violations of this chapter or rules adopted under this chapter may not:

(a) Supervise, be employed by, or manage a shellfish operation licensed or certified under this chapter or rules adopted under this chapter;
(b) Participate in the harvesting, shucking, packing, or shipping of shellfish in commercial quantities or for sale;
(c) Participate in the brokering of shellfish, purchase of shellfish for resale, or retail sale of shellfish; or
(d) Engage in any activity associated with selling or offering to sell shellfish.

(2) Subsections (1)(c) and (d) of this section do not apply to retail purchases of shellfish for personal use.

(3) Subsection (1) of this section applies to a person only during the period of time in which that person’s license or certificate of approval is denied, revoked, or suspended.

(4) Unlawful operations under subsection (1) of this section when a license or certificate of approval is denied, revoked, or suspended is a class C felony. Upon conviction, the department shall order that the person’s license or certificate of approval be revoked for a period of at least five years, or that a person whose application for a license or certificate of approval was denied be ineligible to reapply for a period of at least five years.

(5) A license or certificate of approval issued under this chapter may not be assigned or transferred in any manner without department approval. [2011 c 194 § 7; 1998 c 44 § 1.]

69.30.110 Possession or sale in violation of chapter—Enforcement—Seizure—Disposal. (1) It is unlawful for any person to possess a commercial quantity of shellfish or to sell or offer to sell shellfish in the state which have not been grown, shucked, packed, or shipped in accordance with the provisions of this chapter. Failure of a shellfish grower to display a certificate of approval, or department-approved equivalent, issued under RCW 69.30.050 to an authorized representative of the department, a fish and wildlife officer, or an ex officio fish and wildlife officer subjects the grower to the penalty provisions of this chapter, as well as seizure and disposition, up to and including disposal, of the shellfish by the representative or officer.

(2) Failure of a shellfish processor to display a certificate of approval issued under RCW 69.30.060 to an authorized representative of the department, a fish and wildlife officer, or an ex officio fish and wildlife officer subjects the processor to the penalty provisions of this chapter, as well as seizure and disposition, up to and including disposal, of the shellfish by the representative or officer. [2011 c 194 § 8; 2001 c 253 § 6; 1995 c 147 § 4; 1985 c 51 § 4; 1979 c 141 § 74; 1955 c 144 § 11.]

69.30.120 Inspection by department—Access to regulated business or entity—Administrative inspection warrant. The department may enter and inspect any shellfish growing area or establishment for the purposes of determining compliance with this chapter and rules adopted under this chapter. The department may inspect all shellfish, all permits, all certificates of approval and all records.

During such inspections the department shall have free and unimpeded access to all buildings, yards, warehouses, storage and transportation facilities, vehicles, and other places reasonably considered to be or to have been part of the regulated business or entity, to all ledgers, books, accounts, memorandums, or records required to be compiled or maintained under this chapter or under rules adopted pursuant to this chapter, and to any products, components, or other materials reasonably believed to be or to have been used, processed, or produced by or in connection with the regulated business or activity. In connection with such inspections the department may take such samples or specimens as may be reasonably necessary to determine whether there exists a violation of this chapter or rules adopted under this chapter.

Inspection of establishments may be conducted between eight a.m. and five p.m. on any weekday that is not a legal holiday, during any time the regulated business or entity has established as its usual business hours, at any time the regulated business or entity is open for business or is otherwise in operation, and at any other time with the consent of the owner or authorized agent of the regulated business or entity.

The department may apply for an administrative inspection warrant to a court of competent jurisdiction and an administrative inspection warrant may issue where:
(1) The department has attempted an inspection under this chapter and access to all or part of the regulated business or entity has been actually or constructively denied; or
(2) There is reasonable cause to believe that a violation of this chapter or of rules adopted under this chapter is occurring or has occurred. [1995 c 147 § 5; 1985 c 51 § 5; 1955 c 144 § 12.]

69.30.130 Water pollution laws and rules applicable. All existing laws and rules and regulations governing the pollution of waters of the state shall apply in the control of pollution of shellfish growing areas. [1955 c 144 § 13.]

69.30.140 Penalties. Except as provided in RCW 69.30.085(4), any person convicted of violating any of the provisions of this chapter shall be guilty of a gross misdemeanor. A conviction is an unvacated forfeiture of bail or collateral deposited to secure the defendant’s appearance in court, the payment of a fine, a plea of guilty, or a finding of guilt on a violation of this chapter or rules adopted under this chapter, regardless of whether imposition of sentence is deferred or the penalty is suspended, and shall be treated as a conviction for purposes of license revocation and suspension of privileges under RCW 77.15.700(5). [2011 c 194 § 9; 2001 c 253 § 7; 1995 c 147 § 6; 1985 c 51 § 6; 1955 c 144 § 14.]

*Reviser’s note: RCW 77.15.700 was amended by 2003 c 386 § 2, deleting subsection (5).*

69.30.145 Civil penalties. As limited by RCW 69.30.150, the department may impose civil penalties for violations of standards set forth in this chapter or rules adopted under RCW 69.30.030. [1989 c 200 § 3.]

69.30.150 Civil penalties—General provisions. (1) In addition to any other penalty provided by law, every person who violates standards set forth in this chapter or rules adopted under RCW 69.30.030 is subject to a penalty of not more than five hundred dollars per day for every violation. Every violation is a separate and distinct offense. In case of a continuing violation, every day’s continuance is a separate and distinct violation. Every person who, through an act of commission or omission, procures, aids, or abets in the violation is in violation of this section and is subject to the penalty provided in this section.
(2) The penalty provided for in this section shall be imposed by a notice in writing to the person against whom the civil fine is assessed and shall describe the violation with reasonable particularity. The notice shall be personally served in the manner of service of a summons in a civil action or in a manner which shows proof of receipt. Any penalty imposed by this section shall become due and payable twenty-eight days after notice is received unless application for remission or mitigation is made as provided in subsection (3) of this section or unless application for an adjudicative proceeding is filed as provided in subsection (4) of this section.
(3) Within fourteen days after the notice is received, the person incurring the penalty may apply in writing to the department for the remission or mitigation of the penalty. Upon receipt of the application, the department may remit or mitigate the penalty upon whatever terms the department deems proper, giving consideration to the degree of hazard associated with the violation. The department may only grant a remission or mitigation that it deems to be in the best interests of carrying out the purposes of this chapter. The department may ascertain the facts regarding all such applications in a manner it deems proper. When an application for remission or mitigation is made, any penalty incurred pursuant to this section becomes due and payable twenty-eight days after receipt of the notice setting forth the disposition of the application, unless an application for an adjudicative proceeding to contest the disposition is filed as provided in subsection (4) of this section.
(4) Within twenty-eight days after notice is received, the person incurring the penalty may file an application for an adjudicative proceeding and may pursue subsequent review as provided in chapter 34.05 RCW and applicable rules of the department or board of health.
(5) Any penalty imposed by final order following an adjudicative proceeding becomes due and payable upon service of the final order.
(6) The attorney general may bring an action in the name of the department in the superior court of Thurston county or of any county in which the violator may do business to collect any penalty imposed under this chapter.
(7) All penalties imposed under this section shall be paid to the state treasury and credited to the general fund. [1989 c 200 § 4.]

69.30.900 Severability—1955 c 144. If any provision of this chapter or the application thereof to any person or circumstances shall be held invalid, such invalidity shall not affect the provisions of the application of this chapter which can be given effect without the invalid provision or application, and to this end the provisions of the chapter are declared to be severable. [1955 c 144 § 15.]

Chapter 69.36 RCW

WASHINGTON CAUSTIC POISON ACT OF 1929

Sections

69.36.010 Definitions.
69.36.020 Misbranded sales, etc., prohibited—Exceptions.
69.36.030 Condemnation of misbranded packages.
69.36.040 Enforcement—Approval of labels.
69.36.050 Duty to prosecute.
69.36.060 Penalty.
69.36.070 Short title.

Highway transportation of poisons, corrosives, etc.: RCW 46.48.170, 46.48.175.

69.36.010 Definitions. In this chapter, unless the context or subject matter otherwise requires:
(1) The term "dangerous caustic or corrosive substance" means each and all of the acids, alkalis, and substances named below: (a) Hydrochloric acid and any preparation containing free or chemically unneutralized hydrochloric acid (HCl) in a concentration of ten percent or more; (b) sulphuric acid and any preparation containing free or chemically unneutralized sulphuric acid (H2SO4) in concentration of ten percent or more; (c) nitric acid or any preparation containing free or chemically unneutralized nitric acid (HNO3) in a con-
centration of five percent or more; (d) carboxylic acid (C\textsubscript{6}H\textsubscript{5}O\textsubscript{2}C\textsubscript{2}H\textsubscript{5}OH), otherwise known as phenol, and any preparation containing carboxylic acid in a concentration of five percent or more; (e) oxalic acid and any preparation containing free or chemically unneutralized oxalic acid (H\textsubscript{2}C\textsubscript{2}O\textsubscript{4}) in a concentration of ten percent or more; (f) any salt of oxalic acid and any preparation containing any such salt in a concentration of ten percent or more; (g) acetic acid or any preparation containing free or chemically unneutralized acetic acid (CH\textsubscript{3}COOH) in a concentration of twenty percent or more; (h) hypochlorous acid, either free or combined, and any preparation containing the same in a concentration so as to yield ten percent or more by weight of available chlorine, excluding calx chlorinata, bleaching powder, and chloride of lime; (i) potassium hydroxide and any preparation containing free or chemically unneutralized potassium hydroxide (KOH), including caustic potash and Vienna paste, in a concentration of ten percent or more; (j) sodium hydroxide and any preparation containing free or chemically unneutralized sodium hydroxide (NaOH), including caustic soda and lye, in a concentration of ten percent or more; (k) silver nitrate, sometimes known as lunar caustic, and any preparation containing silver nitrate (AgNO\textsubscript{3}) in a concentration of five percent or more; and (l) ammonia water and any preparation yielding free or chemically uncombined ammonia (NH\textsubscript{3}), including ammonium hydroxide and "hartshorn", in a concentration of five percent or more.

(2) The term "misbranded parcel, package, or container" means a retail parcel, package, or container of any dangerous caustic or corrosive substance for household use, not bearing a conspicuous, easily legible label or sticker, containing (a) the name of the article; (b) the name and place of business of the manufacturer, packer, seller, or distributor; (c) the word "POISON," running parallel with the main body of reading matter on said label or sticker, on a clear, plain background of a distinctly contrasting color, in uncondensed gothic capital letters, the letters to be not less than twenty-four point size, unless there is on said label or sticker no other type so large, in which event the type shall be not smaller than the largest type on the label or sticker; and (d) directions for treatment in case of accidental personal injury by the dangerous caustic or corrosive substance; PROVIDED, That such directions need not appear on labels or stickers on parcels, packages, or containers at the time of shipment or of delivery for shipment by manufacturers or wholesalers for other than household use. PROVIDED FURTHER, That this chapter is not to be construed as applying to any substance subject to the chapter, sold at wholesale or retail for use by a retail druggist in filling prescriptions or in dispensing, in pursuance of a prescription by a physician, dentist, or veterinarian; or for use by or under the direction of a physician, dentist, or veterinarian; or for use by a chemist in the practice or teaching of his or her profession; or for any industrial or professional use, or for use in any of the arts and sciences. [2012 c 117 § 362; 1929 c 82 § 1; RRS § 2508-1. Formerly RCW 69.36.010 and 69.36.020, part.]

### 69.36.020 Misbranded sales, etc., prohibited—Exceptions.

No person shall sell, barter, or exchange, or receive, hold, pack, display, or offer for sale, barter, or exchange, in this state any dangerous caustic or corrosive substance in a misbranded parcel, package, or container, said parcel, package, or container being designed for household use; PROVIDED, That household products for cleaning and washing purposes, subject to this chapter and labeled in accordance therewith, may be sold, offered for sale, held for sale, and distributed in this state by any dealer, wholesale or retail; PROVIDED FURTHER, That no person shall be liable to prosecution and conviction under this chapter when he or she establishes a guaranty bearing the signature and address of a vendor residing in the United States from whom he or she purchased the dangerous caustic or corrosive substance, to the effect that such substance is not misbranded within the meaning of this chapter. No person in this state shall give any such guaranty when such dangerous caustic or corrosive substance is in fact misbranded within the meaning of this chapter. [2012 c 117 § 363; 1929 c 82 § 2; RRS § 2508-2. FORMER PART OF SECTION: 1929 c 82 § 1 now codified in RCW 69.32.010.]

### 69.36.030 Condemnation of misbranded packages.

Any dangerous caustic or corrosive substance in a misbranded parcel, package, or container suitable for household use, that is being sold, bartered, or exchanged, or held, displayed, or offered for sale, barter, or exchange, shall be liable to be proceeded against in any superior court within the jurisdiction of which the same is found and seized for confiscation, and if such substance is condemned as misbranded, by said court, it shall be disposed of by destruction or sale, as the court may direct; and if sold, the proceeds, less the actual costs and charges, shall be paid over to the state treasurer; but such substance shall not be sold contrary to the laws of the state: PROVIDED, HOWEVER, That upon the payment of the costs of such proceedings and the execution and delivery of a good and sufficient bond to the effect that such substance will not be unlawfully sold or otherwise disposed of, the court may by order direct that such substance be delivered to the owner thereof. Such condemnation proceedings shall conform as near as may be to proceedings in the seizure, and condemnation of substances unfit for human consumption. [1929 c 82 § 3; RRS § 2508-3.]

### 69.36.040 Enforcement—Approval of labels.

The director of agriculture shall enforce the provisions of this chapter, and he or she is hereby authorized and empowered to approve and register such brands and labels intended for use under the provisions of this chapter as may be submitted to him or her for that purpose and as may in his or her judgment conform to the requirements of this statute: PROVIDED, HOWEVER, That in any prosecution under this chapter the fact that any brand or label involved in said prosecution has not been submitted to said director for approval, or if submitted, has not been approved by him or her, shall be immaterial. [2012 c 117 § 364; 1929 c 82 § 5; RRS § 2508-5.]

### 69.36.050 Duty to prosecute.

Every prosecuting attorney to whom there is presented, or who in any way procures, satisfactory evidence of any violation of the provisions of this chapter shall cause appropriate proceedings to be commenced and prosecuted in the proper courts, without delay,
for the enforcement of the penalties as in such cases herein
provided. [1929 c 82 § 6; RRS § 2508-6.]

69.38.060 Penalty. Any person violating the provisions of
this chapter shall be guilty of a misdemeanor. [1929 c 82
§ 4; RRS § 2508-4.]

69.38.070 Short title. This chapter may be cited as the
Washington Caustic Poison Act of 1929. [1929 c 82 § 7;
RRS § 2508-7.]

Chapter 69.38 RCW
POISONS—SALES AND MANUFACTURING

69.38.010 "Poison" defined. As used in this chapter
"poison" means:
(1) Arsenic and its preparations;
(2) Cyanide and its preparations, including hydrocyanic
acid;
(3) Strychnine; and
(4) Any other substance designated by the state board of
pharmacy which, when introduced into the human body in
quantities of sixty grains or less, causes violent sickness or
death. [1987 c 34 § 1.]

69.38.020 Exemptions from chapter. All substances
regulated under chapters 15.58, 17.21, 69.04, 69.41, and
69.50 RCW, and chapter 69.45 RCW are exempt from the
provisions of this chapter. [1987 c 34 § 2.]

69.38.030 Poison register—Identification of pur-
chaser. It is unlawful for any person, either on the person’s
own behalf or while an employee of another, to sell any poi-
son without first recording in ink in a "poison register" kept
solely for this purpose the following information:
(1) The date and hour of the sale;
(2) The full name and home address of the purchaser;
(3) The kind and quantity of poison sold; and
(4) The purpose for which the poison is being purchased.

The purchaser shall present to the seller identification
which contains the purchaser’s photograph and signature. No
sale may be made unless the seller is satisfied that the pur-
chaser’s representations are true and that the poison will be
used for a lawful purpose. Both the purchaser and the seller
shall sign the poison register entry.

If a delivery of a poison will be made outside the con-
fines of the seller’s premises, the seller may require the busi-
ness purchasing the poison to submit a letter of authorization
as a substitute for the purchaser’s photograph and signature
requirements. The letter of authorization shall include the
unified business identifier and address of the business, a full
description of how the substance will be used, and the signa-
ture of the purchaser. Either the seller or the employee of the
seller delivering or transferring the poison shall affix his or
her signature to the letter as a witness to the signature and
identification of the purchaser. The transaction shall be
recorded in the poison register as provided in this section.
Letters of authorization shall be kept with the poison register
and shall be subject to the inspection and preservation
requirements contained in RCW 69.38.040. [1988 c 197 § 1;
1987 c 34 § 3.]

69.38.040 Inspection of poison register—Penalty for
failure to maintain register. Every poison register shall be
open for inspection by law enforcement and health officials
at all times and shall be preserved for at least two years after
the date of the last entry. Any person failing to maintain the
poison register as required in this chapter is guilty of a misde-
meanor. [1987 c 34 § 4.]

69.38.050 False representation—Penalty. Any person
making any false representation to a seller when purchasing a
poison is guilty of a gross misdemeanor. [1987 c 34 § 5.]

69.38.060 Manufacturers and sellers of poisons—
License required—Penalty. The state board of pharmacy,
after consulting with the department of health, shall require
and provide for the annual licensure of every person now or
hereafter engaged in manufacturing or selling poisons within
this state. Upon a payment of a fee as set by the department,
the department shall issue a license in such form as it may
prescribe to such manufacturer or seller. Such license shall be
displayed in a conspicuous place in such manufacturer’s or
seller’s place of business for which it is issued.
Any person manufacturing or selling poison within this
state without a license is guilty of a misdemeanor. [1989 1st
ex.s. c 9 § 440; 1987 c 34 § 6.]

Additional notes found at www.leg.wa.gov

Chapter 69.40 RCW
POISONS AND DANGEROUS DRUGS

69.40.010 Poison in edible products. It shall be unlawful
for any person to sell, offer for sale, use, distribute, or
leave in any place, any crackers, biscuit, bread or any other
preparation resembling or in similitude, of any edible prod-
uct, containing arsenic, strychnine or any other poison. [1905
c 141 § 1; RRS § 6140. FORMER PART OF SECTION: 1905
c 141 § 2 now codified as RCW 69.40.015.]

69.40.015 Poison in edible products—Penalty. Any
person violating the provisions of RCW 69.40.010 shall upon
conviction be punished by a fine of not less than ten dollars nor more than five hundred dollars. [1905 c 141 § 2; RRS § 6141. Formerly RCW 69.40.010, part.]

69.40.020 Poison in milk or food products—Penalty. Any person who shall sell, offer to sell, or have in his or her possession for the purpose of sale, either as owner, proprietor, or assistant, or in any manner whatsoever, whether for hire or otherwise, any milk or any food products, containing the chemical ingredient commonly known as formaldehyde, or in which any formaldehyde or other poisonous substance has been mixed, for the purpose of preservation or otherwise, is guilty of a class C felony, and upon conviction thereof shall be imprisoned in the penitentiary for the period of not less than one year nor more than three years. [2003 c 53 § 320; 1905 c 50 § 1; RRS § 6142. Former Part of Section: 1905 c 50 § 2, now codified as RCW 69.40.025.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.40.025 Supplementary to existing laws—Enforcement. *This act shall be supplementary to the laws of this state now in force prohibiting the adulteration of food and fraud in the sale thereof; and the state dairy and food commissioner, the chemist of the state agricultural experiment station, the state attorney general and the prosecuting attorneys of the several counties of this state are hereby required, without additional compensation, to assist in the execution of this act, and in the prosecution of all persons charged with the violation thereof, in like manner and with like powers as they are now authorized and required by law to enforce the laws of this state against the adulteration of food and fraud in the sale thereof. [1905 c 50 § 2; RRS § 6143. Formerly RCW 69.40.020.]

Reviser’s note: *(1) "This act" appears in 1905 c 50 and the sections of the act are codified as RCW 69.40.020 and 69.40.025. (2) The duties of the state dairy and food commissioner have devolved upon the director of agriculture through a chain of statute as follows: 1913 c 60 § 6(2); 1921 c 7 § 93(1). See RCW 43.23.090(1)."

69.40.030 Placing poison or other harmful object or substance in food, drinks, medicine, or water—Penalty. (1) Every person who willfully mingles poison or places any harmful object or substance, including but not limited to pins, tacks, needles, nails, razor blades, wire, or glass in any food, drink, medicine, or other edible substance intended or prepared for the use of a human being or who shall knowingly furnish, with intent to harm another person, any food, drink, medicine, or other edible substance containing such poison or harmful object or substance to another human being, and every person who willfully poisons any spring, well, or reservoir of water, is guilty of a class B felony and shall be punished by imprisonment in a state correctional facility for not less than five years or by a fine of not less than one thousand dollars.

(2) *This act shall not apply to the employer or employees of a person who violates this section without such employer’s knowledge. [2003 c 53 § 321; 1992 c 7 § 48; 1973 c 119 § 1; 1909 c 249 § 264; RRS § 2516. Prior: Code 1881 § 802; 1873 p 185 § 27; 1869 p 202 § 25; 1854 p 79 § 25.]

*Reviser’s note: "this act" refers to the 1973 c 119 § 1 amendment to this section.

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.40.055 Selling repackaged poison without labeling—Penalty. It shall be unlawful for any person to sell at retail or furnish any repackaged poison drug or product without affixing or causing to be affixed to the bottle, box, vessel, or package a label containing the name of the article, all labeling required by the Food and Drug Administration and other federal or state laws or regulations, and the word "poison" distinctly shown with the name and place of the business of the seller.

This section shall not apply to the dispensing of drugs or poisons on the prescription of a practitioner.

The board of pharmacy shall have the authority to promulgate rules for the enforcement and implementation of this section.

Every person who shall violate any of the provisions of this section shall be guilty of a misdemeanor. [1981 c 147 § 4.]

Chapter 69.41 RCW

LEGEND DRUGS—PRESCRIPTION DRUGS

Sections
69.41.010 Definitions.
69.41.020 Prohibited acts—Information not privileged communication.
69.41.030 Sale, delivery, or possession of legend drug without prescription or order prohibited—Exceptions—Penalty.
69.41.040 Prescription requirements—Penalty.
69.41.042 Record requirements.
69.41.044 Confidentiality.
69.41.050 Labeling requirements—Penalty.
69.41.055 Electronic communication of prescription information—Board may adopt rules.
69.41.060 Search and seizure.
69.41.062 Search and seizure at rental premises—Notification of landlord.
69.41.065 Violations—Juvenile driving privileges.
69.41.072 Violations of chapter 69.50 RCW not to be charged under chapter 69.41 RCW—Exception.
69.41.075 Rules—Availability of lists of drugs.
69.41.080 Animal control—Rules for possession and use of legend drugs.
69.41.085 Medication assistance—Community-based care setting.

SUBSTITUTION OF PRESCRIPTION DRUGS

69.41.100 Legislative recognition and declaration.
69.41.110 Definitions.
69.41.120 Prescriptions to contain instruction as to whether or not a therapeutically equivalent generic drug may be substituted—Out-of-state prescriptions—Form—Contents—Procedure.
69.41.130 Savings in price to be passed on to purchaser.
69.41.140 Minimum manufacturing standards and practices.
69.41.150 Liability of practitioner, pharmacist.
69.41.160 Pharmacy signs as to substitution for prescribed drugs.
69.41.170 Coercion of pharmacist prohibited—Penalty.
69.41.180 Rules.
69.41.190 Preferred drug substitution—Exceptions—Notice—Limited restrictions.

IDENTIFICATION OF LEGEND DRUGS—MARKING

69.41.200 Requirements for identification of legend drugs—Marking.
69.41.210 Definitions.
69.41.220 Published lists of drug imprints—Requirements for.
69.41.230 Drugs in violation are contraband.
69.41.240 Rules—Labeling and marking.
69.41.250 Exemptions.
69.41.260 Manufacture or distribution for resale—Requirements.
69.41.280 Confidentiality.
USE OF STEROIDS

69.41.010 Definitions. As used in this chapter, the following terms have the meanings indicated unless the context clearly requires otherwise:

(1) "Administer" means the direct application of a legend drug whether by injection, inhalation, ingestion, or any other means, to the body of a patient or research subject by:
   (a) A practitioner; or
   (b) The patient or research subject at the direction of the practitioner.

(2) "Community-based care settings" include: Community residential programs for the developmentally disabled, certified by the department of social and health services under chapter 71A.12 RCW; adult family homes licensed under chapter 70.128 RCW; and assisted living facilities licensed under chapter 18.20 RCW. Community-based care settings do not include acute care or skilled nursing facilities.

(3) "Deliver" or "delivery" means the actual, constructive, or attempted transfer from one person to another of a legend drug, whether or not there is an agency relationship.

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

(5) "Dispense" means the interpretation of a prescription or order for a legend drug and, pursuant to that prescription or order, the proper selection, measuring, compounding, labeling, or packaging necessary to prepare that prescription or order for delivery.

(6) "Dispenser" means a practitioner who dispenses.

(7) "Distribute" means to deliver other than by administering or dispensing a legend drug.

(8) "Distributor" means a person who distributes.

(9) "Drug" means:
   (a) Substances recognized as drugs in the official United States pharmacopoeia, official homeopathic pharmacopoeia of the United States, or official national formulary, or any supplement to any of them;
   (b) Substances intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in human beings or animals;
   (c) Substances (other than food, minerals or vitamins) intended to affect the structure or any function of the body of human beings or animals; and
   (d) Substances intended for use as a component of any article specified in (a), (b), or (c) of this subsection. It does not include devices or their components, parts, or accessories.

(10) "Electronic communication of prescription information" means the communication of prescription information by computer, or the transmission of an exact visual image of a prescription by facsimile, or other electronic means for original prescription information or prescription refill information for a legend drug between an authorized practitioner and a pharmacy or the transfer of prescription information for a legend drug from one pharmacy to another pharmacy.

(11) "In-home care settings" include an individual's place of temporary and permanent residence, but does not include acute care or skilled nursing facilities, and does not include community-based care settings.

(12) "Legend drugs" means any drugs which are required by state law or regulation of the state board of pharmacy to be dispensed on prescription only or are restricted to use by practitioners only.

(13) "Legible prescription" means a prescription or medication order issued by a practitioner that is capable of being read and understood by the pharmacist filling the prescription or the nurse or other practitioner implementing the medication order. A prescription must be hand printed, typewritten, or electronically generated.

(14) "Medication assistance" means assistance rendered by a nonpractitioner to an individual residing in a community-based care setting or in-home care setting to facilitate the individual's self-administration of a legend drug or controlled substance. It includes reminding or coaching the individual, handing the medication container to the individual, opening the individual's medication container, using an enabler, or placing the medication in the individual's hand, and such other means of medication assistance as defined by rule adopted by the department. A nonpractitioner may help in the preparation of legend drugs or controlled substances for self-administration where a practitioner has determined and communicated orally or by written direction that such medication assistance is necessary and appropriate. Medication assistance shall not include assistance with intravenous medications or injectable medications, except prefilled insulin syringes.

(15) "Person" means individual, corporation, government or governmental subdivision or agency, business trust, estate, trust, partnership or association, or any other legal entity.

(16) "Practitioner" means:
   (a) A physician under chapter 18.71 RCW, an osteopathic physician or an osteopathic physician and surgeon under chapter 18.57 RCW, a dentist under chapter 18.32 RCW, a podiatric physician and surgeon under chapter 18.22 RCW, a veterinarian under chapter 18.92 RCW, a registered nurse, advanced registered nurse practitioner, or licensed practical nurse under chapter 18.79 RCW, an optometrist under chapter 18.53 RCW who is certified by the optometry board under RCW 18.53.010, an osteopathic physician assistant under chapter 18.57A RCW, a physician assistant under chapter 18.71A RCW, a naturopath licensed under chapter 18.36A RCW, a pharmacist under chapter 18.64 RCW, or, when acting under the required supervision of a dentist licensed under chapter 18.32 RCW, a dental hygienist licensed under chapter 18.29 RCW;
   (b) A pharmacy, hospital, or other institution licensed, registered, or otherwise permitted to distribute, dispense, conduct research with respect to, or to administer a legend drug in the course of professional practice or research in this state; and
   (c) A physician licensed to practice medicine and surgery or a physician licensed to practice osteopathic medicine and surgery in any state, or province of Canada, which shares a common border with the state of Washington.
(17) "Secretary" means the secretary of health or the secretary's designee. [2012 c 10 § 44; 2009 c 549 § 1024; 2006 c 8 § 115. Prior: 2003 c 257 § 2; 2003 c 140 § 11; 2000 c 8 § 2; prior: 1998 c 222 § 1; 1998 c 70 § 2; 1996 c 178 § 16; 1994 sps. c 9 § 736; prior: 1989 1st ex.s. c 9 § 426; 1989 c 36 § 3; 1984 c 153 § 17; 1980 c 71 § 1; 1979 ex.s. c 139 § 1; 1973 1st ex.s. c 186 § 1.]

Application—2012 c 10: See note following RCW 18.20.010.

Findings—2006 c 8: "The legislature finds that prescription drug errors occur because the pharmacist or nurse cannot read the prescription from the physician or other provider with prescriptive authority. The legislature further finds that legible prescriptions can prevent these errors." [2006 c 8 § 114.]

Findings—Intent—Part headings and subheadings not law—Severability—2006 c 8: See notes following RCW 5.64.010.

Effective date—2003 c 140: See note following RCW 18.79.040.

Findings—Intent—2000 c 8: "The legislature finds that we have one of the finest health care systems in the world and excellent professionals to deliver that care. However, there are incidents of medication errors that are avoidable and serious mistakes that are preventable. Medical errors throughout the health care system constitute one of the nation’s leading causes of death and injury resulting in over seven thousand deaths a year, according to a recent report from the institute of medicine. The majority of medical errors do not result from individual recklessness, but from basic flaws in the way the health system is organized. There is a need for a comprehensive strategy for government, industry, consumers, and health providers to reduce medical errors. The legislature declares a need to bring about greater safety for patients in this state who depend on prescription drugs.

It is the intent of the legislature to promote medical safety as a top priority for all citizens of our state." [2000 c 8 § 1.]

Additional notes found at www.leg.wa.gov

69.41.020 Prohibited acts—Information not privileged communication. Legend drugs shall not be sold, delivered, dispensed or administered except in accordance with this chapter.

(1) No person shall obtain or attempt to obtain a legend drug, or procure or attempt to procure the administration of a legend drug:

(a) By fraud, deceit, misrepresentation, or subterfuge; or

(b) By the forgery or alteration of a prescription or of any written order; or

(c) By the concealment of a material fact; or

(d) By the use of a false name or the giving of a false address.

(2) Information communicated to a practitioner in an effort unlawfully to procure a legend drug, or unlawfully to procure the administration of any such drug, shall not be deemed a privileged communication.

(3) No person shall willfully make a false statement in any prescription, order, report, or record, required by this chapter.

(4) No person shall, for the purpose of obtaining a legend drug, falsely assume the title of, or represent himself or herself to be, a manufacturer, wholesaler, or any practitioner.

(5) No person shall make or utter any false or forged prescription or other written order for legend drugs.

(6) No person shall affix any false or forged label to a package or receptacle containing legend drugs.

(7) No person shall willfully fail to maintain the records required by RCW 69.41.042 and *69.41.270.

(8) A violation of this section is a class B felony punishable according to chapter 9A.20 RCW. [2003 c 53 § 322. Prior: 1989 1st ex.s. c 9 § 408; 1989 c 352 § 8; 1973 1st ex.s. c 186 § 2.]

Reviser's note: RCW 69.41.270 was repealed by 2003 c 275 § 5.

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

Additional notes found at www.leg.wa.gov

69.41.030 Sale, delivery, or possession of legend drug without prescription or order prohibited—Exceptions—Penalty. (1) It shall be unlawful for any person to sell, deliver, or possess any legend drug except upon the order or prescription of a physician under chapter 18.71 RCW, an osteopathic physician and surgeon under chapter 18.57 RCW, an optometrist licensed under chapter 18.53 RCW who is certified by the optometry board under RCW 18.53.010, a dentist under chapter 18.32 RCW, a podiatric physician and surgeon under chapter 18.22 RCW, a veterinarian under chapter 18.92 RCW, a commissioned medical or dental officer in the United States armed forces or public health service in the discharge of his or her official duties, a duly licensed physician or dentist employed by the veterans administration in the discharge of his or her official duties, a registered nurse or advanced registered nurse practitioner under chapter 18.79 RCW when authorized by the nursing care quality assurance commission, an osteopathic physician assistant under chapter 18.57A RCW when authorized by the board of osteopathic medicine and surgery, a physician assistant under chapter 18.71A RCW when authorized by the medical quality assurance commission, or any of the following professionals in any province of Canada that shares a common border with the state of Washington or in any state of the United States: A physician licensed to practice medicine and surgery or a physician licensed to practice osteopathic medicine and surgery, a dentist licensed to practice dentistry, a podiatric physician and surgeon licensed to practice podiatric medicine and surgery, a licensed advanced registered nurse practitioner, or a veterinarian licensed to practice veterinary medicine: PROVIDED, HOWEVER, That the above provisions shall not apply to sale, delivery, or possession by drug wholesalers or drug manufacturers, or their agents or employees, or to any practitioner acting within the scope of his or her license, or to a common or contract carrier or warehouse operator, or any employee thereof, whose possession of any legend drug is in the usual course of business or employment: PROVIDED FURTHER, That nothing in this chapter or chapter 18.64 RCW shall prevent a family planning clinic that is under contract with the health care authority from selling, delivering, possessing, and dispensing commercially prepackaged oral contraceptives prescribed by authorized, licensed health care practitioners.

(2)(a) A violation of this section involving the sale, delivery, or possession with intent to sell or deliver is a class B felony punishable according to chapter 9A.20 RCW.

(b) A violation of this section involving possession is a misdemeanor. [2011 1st sps. c 15 § 79; 2011 c 336 § 837; 2010 c 83 § 1. Prior: 2003 c 142 § 3; 2003 c 53 § 323; 1996 c 178 § 17; 1994 sps. c 9 § 737; 1991 c 30 § 1; 1990 c 219 § 2; 1987 c 144 § 1; 1981 c 120 § 1; 1979 ex.s. c 139 § 2; 1977 c 69 § 1; 1973 1st ex.s. c 186 § 3.]

Reviser's note: This section was amended by 2011 c 336 § 837 and by 2011 1st sps. c 15 § 79, each without reference to the other. Both amend-

Severability—2003 c 142: See note following RCW 18.53.010.

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

Finding—1990 c 219: "The legislature finds that Washington citizens in the border areas of this state are prohibited from having prescriptions from out-of-state dentists and veterinarians filled at their in-state pharmacies, and that it is in the public interest to remove this barrier for the state’s citizens."
[1990 c 219 § 1.]

Additional notes found at www.leg.wa.gov

69.41.032 Prescription of legend drugs by dialysis programs. This chapter shall not prevent a medicare-approved dialysis center or facility operating a medicare-approved home dialysis program from selling, delivering, possessing, or dispensing directly to its dialysis patients, in case or full shelf lots, if prescribed by a physician licensed under chapter 18.57 or 18.71 RCW, those legend drugs determined by the board pursuant to rule. [1987 c 41 § 2.]

Application of pharmacy statutes to dialysis programs: RCW 18.64.257.

69.41.040 Prescription requirements—Penalty. (1) A prescription, in order to be effective in legalizing the possession of legend drugs, must be issued for a legitimate medical purpose by one authorized to prescribe the use of such legend drugs. An order purporting to be a prescription issued to a drug abuser or habitual user of legend drugs, not in the course of professional treatment, is not a prescription within the meaning and intent of this section; and the person who possesses, or dispensing directly to its dialysis patients, in case or full shelf lots, if prescribed by a physician licensed under chapter 18.57 or 18.71 RCW, those legend drugs determined by the board pursuant to rule. [1987 c 41 § 2.]

Application of pharmacy statutes to dialysis programs: RCW 18.64.257.

69.41.055 Electronic communication of prescription information—Board may adopt rules. (1) Information concerning an original prescription or information concerning a prescription refill for a legend drug may be electronically communicated between an authorized practitioner and a pharmacy of the patient’s choice with no intervening person having access to the prescription drug order pursuant to the provisions of this chapter if the electronically communicated prescription information complies with the following:

(a) Electronically communicated prescription information must comply with all applicable statutes and rules regarding the form, content, recordkeeping, and processing of a prescription for a legend drug;

(b) The system used for transmitting electronically communicated prescription information and the system used for receiving electronically communicated prescription information must be approved by the board. This subsection does not apply to currently used facsimile equipment transmitting an exact visual image of the prescription. The board shall maintain and provide, upon request, a list of systems used for electronically communicating prescription information currently approved by the board;

(c) An explicit opportunity for practitioners must be made to indicate their preference on whether a therapeutically equivalent generic drug may be substituted;

(d) Prescription drug orders are confidential health information, and may be released only to the patient or the patient’s authorized representative, the prescriber or other authorized practitioner then caring for the patient, or other persons specifically authorized by law to receive such information;

(e) To maintain confidentiality of prescription records, the electronic system shall have adequate security and systems safeguards designed to prevent and detect unauthorized
access, modification, or manipulation of these records. The pharmacist in charge shall establish or verify the existence of policies and procedures which ensure the integrity and confidentiality of prescription information transmitted to the pharmacy by electronic means. All managers, employees, and agents of the pharmacy are required to read, sign, and comply with the established policies and procedures; and

(1) The pharmacist shall exercise professional judgment regarding the accuracy, validity, and authenticity of the prescription drug order received by way of electronic transmission, consistent with federal and state laws and rules and guidelines of the board.

(2) The board may adopt rules implementing this section. [1998 c 222 § 2.]

69.41.060 Search and seizure. If, upon the sworn complaint of any person, it shall be made to appear to any judge of the superior or district court that there is probable cause to believe that any legend drug is being used, manufactured, sold, bartered, exchanged, given away, furnished or otherwise disposed of or kept in violation of the provisions of this chapter, such judge shall, with or without the approval of the prosecuting attorney, issue a warrant directed to any peace officer in the county, commanding the peace officer to search the premises designated and described in such complaint and warrant, and to seize all legend drugs there found, together with the vessels in which they are contained, and all implements, furniture and fixtures used or kept for the illegal manufacture, sale, barter, exchange, giving away, furnishing or otherwise disposing of such legend drugs and to safely keep the same, and to make a return of said warrant within three days, showing all acts and things done thereunder, with a particular statement of all articles seized and the name of the person or persons in whose possession the same were found, if any, and if no person be found in the possession of said articles, the returns shall so state. A copy of said warrant shall be served upon the person or persons found in possession of any such legend drugs, furniture or fixtures so seized, and if no person be found in the possession thereof, a copy of said warrant shall be posted on the door of the building or room wherein the same are found, or, if there be no door, then in any conspicuous place upon the premises. [1987 c 202 § 227; 1973 1st ex.s. c 186 § 6.]

Intent—1987 c 202: See note following RCW 2.04.190.

69.41.062 Search and seizure at rental premises—Notification of landlord. Whenever a legend drug which is sold, delivered, or possessed in violation of this chapter is seized at rental premises, the law enforcement agency shall make a reasonable attempt to discover the identity of the landlord and shall notify the landlord in writing, at the last address listed in the property tax records and at any other address known by the law enforcement agency, of the seizure and the location of the seizure. [1988 c 150 § 8.]

Legislative findings—Severability—1988 c 150: See notes following RCW 59.18.130.

69.41.065 Violations—Juvenile driving privileges. (1) If a juvenile thirteen years of age or older and under the age of twenty-one is found by a court to have committed any offense that is a violation of this chapter, the court shall notify the department of licensing within twenty-four hours after entry of the judgment.

(2) Except as otherwise provided in subsection (3) of this section, upon petition of a juvenile whose privilege to drive has been revoked pursuant to RCW 46.20.265, the court may notify the department of licensing that the juvenile’s privilege to drive should be reinstated.

(3) If the conviction is for the juvenile’s first violation of this chapter or chapter 66.44, 69.50, or 69.52 RCW, the juvenile may not petition the court for reinstatement of the juvenile’s privilege to drive revoked pursuant to RCW 46.20.265 until the later of ninety days after the date the juvenile turns sixteen or ninety days after the judgment was entered. If the conviction was for the juvenile’s second or subsequent violation of this chapter or chapter 66.44, 69.50, or 69.52 RCW, the juvenile may not petition the court for reinstatement of the juvenile’s privilege to drive revoked pursuant to RCW 46.20.265 until the later of the date the juvenile turns seventeen or one year after the date judgment was entered. [1989 c 271 § 119; 1988 c 148 § 4.]

Legislative finding—Severability—1988 c 148: See notes following RCW 13.40.265.

Additional notes found at www.leg.wa.gov

69.41.072 Violations of chapter 69.50 RCW not to be charged under chapter 69.41 RCW—Exception. Any offense which is a violation of chapter 69.50 RCW other than RCW 69.50.4012 shall not be charged under this chapter. [2003 c 53 § 327.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.41.075 Rules—Availability of lists of drugs. The state board of pharmacy may make such rules for the enforcement of this chapter as are deemed necessary or advisable. The board shall identify, by rule-making pursuant to chapter 34.05 RCW, those drugs which may be dispensed only on prescription or are restricted to use by practitioners, only. In so doing the board shall consider the toxicity or other potentiality for harmful effect of the drug, the method of its use, and any collateral safeguards necessary to its use. The board shall classify a drug as a legend drug where these considerations indicate the drug is not safe for use except under the supervision of a practitioner.

In identifying legend drugs the board may incorporate in its rules lists of drugs contained in commercial pharmaceutical publications by making specific reference to each such list and the date and edition of the commercial publication containing it. Any such lists so incorporated shall be available for public inspection at the headquarters of the department of health and shall be available on request from the department of health upon payment of a reasonable fee to be set by the department. [1989 1st ex.s. c 9 § 427; 1979 ex.s. c 139 § 3.]

Additional notes found at www.leg.wa.gov

69.41.080 Animal control—Rules for possession and use of legend drugs. Humane societies and animal control agencies registered with the state board of pharmacy under chapter 69.50 RCW and authorized to euthanize animals may purchase, possess, and administer approved legend drugs for use of animals.
the sole purpose of sedating animals prior to euthanasia, when necessary, and for use in chemical capture programs. For the purposes of this section, "approved legend drugs" means those legend drugs designated by the board by rule as being approved for use by such societies and agencies for animal sedating or capture and does not include any substance regulated under chapter 69.50 RCW. Any society or agency so registered shall not permit persons to administer any legend drugs unless such person has demonstrated to the satisfaction of the board adequate knowledge of the potential hazards involved in and the proper techniques to be used in administering the drugs.

The board shall promulgate rules to regulate the purchase, possession, and administration of legend drugs by such societies and agencies and to insure strict compliance with the provisions of this section. Such rules shall require that the storage, inventory control, administration, and recordkeeping for approved legend drugs conform to the standards adopted by the board under chapter 69.50 RCW to regulate the use of controlled substances by such societies and agencies. The board may suspend or revoke a registration under chapter 69.50 RCW upon a determination by the board that the person administering legend drugs has not demonstrated adequate knowledge as herein provided. This authority is granted in addition to any other power to suspend or revoke a registration as provided by law. [1989 c 242 § 1.]

69.41.085 Medication assistance—Community-based care setting. Individuals residing in community-based care settings, such as adult family homes, assisted living facilities, and residential care settings for individuals with developmental disabilities, including an individual’s home, may receive medication assistance. Nothing in this chapter affects the right of an individual to refuse medication or requirements relating to informed consent. [2012 c 10 § 45; 2003 c 140 § 12; 1998 c 70 § 1.]

Application—2012 c 10: See note following RCW 18.20.010.
Effective date—2003 c 140: See note following RCW 18.79.040.

SUBSTITUTION OF PRESCRIPTION DRUGS

69.41.100 Legislative recognition and declaration. The legislature recognizes the responsibility of the state to insure that the citizens of the state are offered a choice between generic drugs and brand name drugs and the benefit of quality pharmaceutical products at competitive prices. Advances in the drug industry resulting from research and the elimination of counterfeiting of prescription drugs should benefit the users of the drugs. Pharmacy must continue to operate with accountability and effectiveness. The legislature hereby declares it to be the policy of the state that its citizens receive safe and therapeutically effective drug products at the most reasonable cost consistent with high drug quality standards. [1986 c 52 § 1; 1977 ex.s. c 352 § 1.]

Additional notes found at www.leg.wa.gov

69.41.110 Definitions. As used in RCW 69.41.100 through 69.41.180, the following words shall have the following meanings:

1. "Brand name" means the proprietary or trade name selected by the manufacturer and placed upon a drug, its container, label, or wrapping at the time of packaging;
2. "Generic name" means the official title of a drug or ingredient published in the latest edition of a nationally recognized pharmacopoeia or formulary;
3. "Substitute" means to dispense, with the practitioner’s authorization, a "therapeutically equivalent" drug product of the identical base or salt as the specific drug product prescribed: PROVIDED, That with the practitioner’s prior consent, therapeutically equivalent drugs other than the identical base or salt may be dispensed;
4. "Therapeutically equivalent" means essentially the same efficacy and toxicity when administered in an individual in the same dosage regimen; and
5. "Practitioner" means a physician, osteopathic physician and surgeon, dentist, veterinarian, or any other person authorized to prescribe drugs under the laws of this state. [1979 c 110 § 1; 1977 ex.s. c 352 § 2.]

69.41.120 Prescriptions to contain instruction as to whether or not a therapeutically equivalent generic drug may be substituted—Out-of-state prescriptions—Form—Contents—Procedure. Every drug prescription shall contain an instruction on whether or not a therapeutically equivalent generic drug may be substituted in its place, unless substitution is permitted under a prior-consent authorization.

If a written prescription is involved, the prescription must be legible and the form shall have two signature lines at opposite ends on the bottom of the form. Under the line at the right side shall be clearly printed the words "DISPENSE AS WRITTEN". Under the line at the left side shall be clearly printed the words "SUBSTITUTION PERMITTED". The practitioner shall communicate the instructions to the pharmacist by signing the appropriate line. No prescription shall be valid without the signature of the practitioner on one of these lines. In the case of a prescription issued by a practitioner in another state that uses a one-line prescription form or variation thereof, the pharmacist may substitute a therapeutically equivalent generic drug unless otherwise instructed by the practitioner through the use of the words "dispense as written", words of similar meaning, or some other indication.

If an oral prescription is involved, the practitioner or the practitioner’s agent shall instruct the pharmacist as to whether or not a therapeutically equivalent generic drug may be substituted in its place. The pharmacist shall note the instructions on the file copy of the prescription.

The pharmacist shall note the manufacturer of the drug dispensed on the file copy of a written or oral prescription. [2000 c 8 § 3; 1990 c 218 § 1; 1979 c 110 § 2; 1977 ex.s. c 352 § 3.]

Findings—Intent—2000 c 8: See note following RCW 69.41.010.

69.41.130 Savings in price to be passed on to purchaser. Unless the brand name drug is requested by the patient or the patient’s representative, the pharmacist shall substitute an equivalent drug product which he or she has in stock if its wholesale price to the pharmacist is less than the wholesale price of the prescribed drug product, and at least sixty percent of the savings shall be passed on to the pur-
69.41.140 Minimum manufacturing standards and practices. A pharmacist may not substitute a product under the provisions of this section unless the manufacturer has shown that the drug has been manufactured with the following minimum good manufacturing standards and practices:

(1) Maintain quality control standards equal to those of the Food and Drug Administration;

(2) Comply with regulations promulgated by the Food and Drug Administration. [1979 c 110 § 4; 1977 ex.s. c 352 § 5.]

69.41.150 Liability of practitioner, pharmacist. (1) A practitioner who authorizes a prescribed drug shall not be liable for any side effects or adverse reactions caused by the manner or method by which a substituted drug product is selected or dispensed.

(2) A pharmacist who substitutes an equivalent drug product pursuant to RCW 69.41.100 through 69.41.180 as now or hereafter amended assumes no greater liability for selecting the dispensed drug product than would be incurred in filling a prescription for a drug product prescribed by its established name.

(3) A pharmacist who substitutes a preferred drug for a nonpreferred drug pursuant to RCW 69.41.190 assumes no greater liability for substituting the preferred drug than would be incurred in filling a prescription for the preferred drug when prescribed by name. [2003 1st sp.s. c 29 § 6; 1979 c 110 § 5; 1977 ex.s. c 352 § 6.]

Finding—Intent—Severability—Conflict with federal requirements—Effective date—2003 1st sp.s. c 29: See notes following RCW 74.09.650.

69.41.160 Pharmacy signs as to substitution for prescribed drugs. Every pharmacy shall post a sign at a location at the prescription counter that is readily visible to patrons stating, "Under Washington law, an equivalent but less expensive drug may in some cases be substituted for the drug prescribed by your doctor. Such substitution, however, may only be made with the consent of your doctor. Please consult your pharmacist or physician for more information." [1979 c 110 § 6; 1977 ex.s. c 352 § 7.]

69.41.170 Coercion of pharmacist prohibited—Penalty. It shall be unlawful for any employer to coerce, within the meaning of RCW 9A.36.070, any pharmacist to dispense a generic drug or to substitute a generic drug for another drug. A violation of this section shall be punishable as a misdemeanor. [1977 ex.s. c 352 § 8.]

69.41.180 Rules. The state board of pharmacy may adopt any necessary rules under chapter 34.05 RCW for the implementation, continuation, or enforcement of RCW 69.41.100 through 69.41.180, including, but not limited to, a list of therapeutically or nontherapeutically equivalent drugs which, when adopted, shall be provided to all registered pharmacists in the state and shall be updated as necessary. [1979 c 110 § 7; 1977 ex.s. c 352 § 9.]

69.41.190 Preferred drug substitution—Exceptions—Notice—Limited restrictions. (1)(a) Except as provided in subsection (2) of this section, any pharmacist filling a prescription under a state purchased health care program as defined in *RCW 41.05.011(2) shall substitute, where identified, a preferred drug for any nonpreferred drug in a given therapeutic class, unless the endorsing practitioner has indicated on the prescription that the nonpreferred drug must be dispensed as written, or the prescription is for a refill of an antipsychotic, antidepressant, antiepileptic, chemotherapy, antiretroviral, or immunosuppressive drug, or for the refill of a immunomodulator/antiviral treatment for hepatitis C for which an established, fixed duration of therapy is prescribed for at least twenty-four weeks but no more than forty-eight weeks, in which case the pharmacist shall dispense the prescribed nonpreferred drug.

(b) When a substitution is made under (a) of this subsection, the dispensing pharmacist shall notify the prescribing practitioner of the specific drug and dose dispensed.

(2)(a) A state purchased health care program may impose limited restrictions on an endorsing practitioner’s authority to write a prescription to dispense as written only under the following circumstances:

(i) There is statistical or clear data demonstrating the endorsing practitioner’s frequency of prescribing dispensed as written for nonpreferred drugs varies significantly from the prescribing patterns of his or her peers;

(ii) The medical director of a state purchased health program has: (A) Presented the endorsing practitioner with data that indicates the endorsing practitioner’s prescribing patterns vary significantly from his or her peers, (B) provided the endorsing practitioner an opportunity to explain the variation in his or her prescribing patterns to those of his or her peers, and (C) if the variation in prescribing patterns cannot be explained, provided the endorsing practitioner sufficient time to change his or her prescribing patterns to align with those of his or her peers; and

(iii) The restrictions imposed under (a) of this subsection (2) must be limited to the extent possible to reduce variation in prescribing patterns and shall remain in effect only until such time as the endorsing practitioner can demonstrate a reduction in variation in line with his or her peers.

(b) A state purchased health care program may immediately designate an available, less expensive, equally effective generic product in a previously reviewed drug class as a preferred drug, without first submitting the product to review by the pharmacy and therapeutics committee established pursuant to RCW 70.14.050.

(c) For a patient’s first course of treatment within a therapeutic class of drugs, a state purchased health care program may impose limited restrictions on endorsing practitioners’ authority to write a prescription to dispense as written, only under the following circumstances:

(i) There is a less expensive, equally effective therapeutic alternative generic product available to treat the condition;

(ii) The drug use review board established under WAC 388-530-4000 reviews and provides recommendations as to the appropriateness of the limitation;

(iii) Notwithstanding the limitation set forth in (c)(ii) of this subsection (2), the endorsing practitioner shall have an
opportunity to request as medically necessary, that the brand name drug be prescribed as the first course of treatment;

(iv) The state purchased health care program may provide, where available, prescription, emergency room, diagnosis, and hospitalization history with the endorsing practitioner; and

(v) Specifically for antipsychotic restrictions, the state purchased health care program shall effectively guide good practice without interfering with the timeliness of clinical decision making. Health care authority prior authorization programs must provide for responses within twenty-four hours and at least a seventy-two hour emergency supply of the requested drug.

(d) If, within a therapeutic class, there is an equally effective therapeutic alternative over-the-counter drug available, a state purchased health care program may designate the over-the-counter drug as the preferred drug.

(e) A state purchased health care program may impose limited restrictions on endorsing practitioners’ authority to prescribe pharmaceuticals to be dispensed as written for a purpose outside the scope of their approved labels only under the following circumstances:

(i) There is a less expensive, equally effective on-label product available to treat the condition;

(ii) The drug use review board established under WAC 388-530-4000 reviews and provides recommendations as to the appropriateness of the limitation; and

(iii) Notwithstanding the limitation set forth in (e)(ii) of this subsection (2), the endorsing practitioner shall have an opportunity to request as medically necessary, that the drug be prescribed for a covered off-label purpose.

(f) The provisions of this subsection related to the definition of medically necessary, prior authorization procedures and patient appeal rights shall be implemented in a manner consistent with applicable federal and state law.

(3) Notwithstanding the limitations in subsection (2) of this section, for refills for an antipsychotic, antidepressant, antiepileptic, chemotherapy, antiretroviral, or immunosuppressive drug, or for the refill of an immunomodulator antiviral treatment for hepatitis C for which an established, fixed duration of therapy is prescribed for at least twenty-four weeks by no more than forty-eight weeks, the pharmacist shall dispense the prescribed nonpreferred drug. [2011 1st sp.s. c 15 § 80; 2009 c 575 § 1; 2006 c 233 § 1; 2003 1st sp.s. c 29 § 5.]

*Reviser’s note: RCW 41.05.011 was alphabetized pursuant to RCW 1.08.015(2)(k), changing subsection (2) to subsection (21).*

**Effective date—Findings—Intent—Report—Agency transfer—References to head of health care authority—Draft legislation—2011 1st sp.s. c 15:** See notes following RCW 74.09.010.

**Effective date—2009 c 575:** "This act is necessary for the immediate preservation of the public peace, health, or safety, or support of the state government and its existing public institutions, and takes effect immediately [May 19, 2009]." [2009 c 575 § 2.]

**Finding—Intent—Severability—Conflict with federal requirements—Effective date—2003 1st sp.s. c 29:** See notes following RCW 74.09.650.

**IDENTIFICATION OF LEGEND DRUGS—MARKING**

### 69.41.200 Requirements for identification of legend drugs—Marking.

(1) No legend drug in solid dosage form may be manufactured or commercially distributed within this state unless it has clearly marked or imprinted on it an individual symbol, number, company name, words, letters, marking, or National Drug Code number identifying the drug and the manufacturer or distributor of such drug.

(2) No manufacturer or distributor may sell any legend drug contained within a bottle, vial, carton, or other container, or in any way affixed or appended to or enclosed within a package of any kind designed or intended for delivery in such container or package to an ultimate consumer within this state unless such container or package has clearly and permanently marked or imprinted on it an individual symbol, number, company name, words, letters, marking, or National Drug Code number identifying the drug and the manufacturer or distributor of such drug.

(3) Whenever the distributor of a legend drug does not also manufacture it, the names and places of businesses of both shall appear on the stock container or package label in words that truly distinguish each. [1980 c 83 § 1.]

### 69.41.210 Definitions.

The terms defined in this section shall have the meanings indicated when used in RCW 69.41.200 through 69.41.260.

(1) "Distributor" means any corporation, person, or other entity which distributes for sale a legend drug under its own label even though it is not the actual manufacturer of the legend drug.

(2) "Solid dosage form" means capsules or tablets or similar legend drug products intended for administration and which could be ingested orally.

(3) "Legend drug" means any drugs which are required by state law or regulation of the board to be dispensed as prescription only or are restricted to use by prescribing practitioners only and shall include controlled substances in Schedules I through V of chapter 69.50 RCW.

(4) "Board" means the state board of pharmacy. [1980 c 83 § 2.]

### 69.41.220 Published lists of drug imprints—Requirements for.

Each manufacturer and distributor shall publish and provide to the board by filing with the department printed material which will identify each current imprint used by the manufacturer or distributor. The board shall be notified of any change by the filing of any change with the department. This information shall be provided by the department to all pharmacies licensed in the state of Washington, poison control centers, and hospital emergency rooms. [1989 1st ex.s. c 9 § 428; 1980 c 83 § 3.]

Additional notes found at www.leg.wa.gov

### 69.41.230 Drugs in violation are contraband.

Any legend drug prepared or manufactured or offered for sale in violation of this chapter or implementing rules shall be contraband and subject to seizure under the provisions of RCW 69.41.060. [1980 c 83 § 4.]

### 69.41.240 Rules—Labeling and marking.

The board shall have authority to promulgate rules and regulations for the enforcement and implementation of RCW 69.41.050 and 69.41.200 through 69.41.260. [1980 c 83 § 5.]
69.41.250 Exemptions. (1) The board, upon application of a manufacturer, may exempt a particular legend drug from the requirements of RCW 69.41.050 and 69.41.200 through 69.41.260 on the grounds that imprinting is infeasible because of size, texture, or other unique characteristics.

(2) The provisions of RCW 69.41.050 and 69.41.200 through 69.41.260 shall not apply to any legend drug which is prepared or manufactured by a pharmacy in this state and is for the purpose of retail sale from such pharmacy and not intended for resale. [1980 c 83 § 6.]

69.41.260 Manufacture or distribution for resale—Requirements. All legend drugs manufactured or distributed for resale to any entity in this state other than the ultimate consumer shall meet the requirements of RCW 69.41.050 and 69.41.200 through 69.41.260 from a date eighteen months after June 12, 1980. [1980 c 83 § 7.]

69.41.280 Confidentiality. All records, reports, and information obtained by the board or its authorized representatives from or on behalf of a pharmaceutical manufacturer, representative of a manufacturer, wholesaler, pharmacy, or practitioner who purchases, dispenses, or distributes legend drugs under this chapter are confidential and exempt from public inspection and copying under chapter 42.56 RCW. Nothing in this section restricts the investigations or the proceedings of the board so long as the board and its authorized representatives comply with the provisions of chapter 42.56 RCW. [2005 c 274 § 329; 1989 c 352 § 6.]

Part headings not law—Effective date—2005 c 274: See RCW 42.56.901 and 42.56.902.

USE OF STEROIDS

69.41.300 Definitions. For the purposes of RCW 69.41.300 through 69.41.350, "steroids" shall include the following:

(1) "Anabolic steroids" means synthetic derivatives of testosterone or any isomer, ester, salt, or derivative that act in the same manner on the human body;

(2) "Androgens" means testosterone in one of its forms or a derivative, isomer, ester, or salt, that act in the same manner on the human body; and

(3) "Human growth hormones" means growth hormones, or a derivative, isomer, ester, or salt that act in the same manner on the human body. [2003 c 53 § 328; 1989 c 369 § 1.]

Intent—Effective date—2003 c 53: See notes following RCW 42.48.180.

69.41.310 Rules. The state board of pharmacy shall specify by rule drugs to be classified as steroids as defined in RCW 69.41.300.

On or before December 1 of each year, the board shall inform the appropriate legislative committees of reference of the drugs that the board has added to the steroids in RCW 69.41.300. The board shall submit a statement of rationale for the changes. [1989 c 369 § 2.]

69.41.320 Practitioners—Restricted use—Medical records. (1)(a) A practitioner shall not prescribe, administer, or dispense steroids, as defined in RCW 69.41.300, or any form of autotransfusion for the purpose of manipulating hormones to increase muscle mass, strength, or weight, or for the purpose of enhancing athletic ability, without a medical necessity to do so.

(b) A person violating this subsection is guilty of a gross misdemeanor and is subject to disciplinary action under RCW 18.130.180.

(2) A practitioner shall complete and maintain patient medical records which accurately reflect the prescribing, administering, or dispensing of any substance or drug described in this section or any form of autotransfusion. Patient medical records shall indicate the diagnosis and purpose for which the substance, drug, or autotransfusion is prescribed, administered, or dispensed and any additional information upon which the diagnosis is based. [2003 c 53 § 329; 1989 c 369 § 3.]

Intent—Effective date—2003 c 53: See notes following RCW 42.48.180.

69.41.330 Public warnings—School districts. The superintendent of public instruction shall develop and distribute to all school districts signs of appropriate design and dimensions advising students of the health risks that steroids present when used solely to enhance athletic ability, and of the penalties for their unlawful possession provided by RCW 69.41.300 through 69.41.350.

School districts shall post or cause the signs to be posted in a prominent place for ease of viewing on the premises of school athletic departments. [2003 c 53 § 330; 1989 c 369 § 5.]

Intent—Effective date—2003 c 53: See notes following RCW 42.48.180.

69.41.340 Student athletes—Violations—Penalty. The superintendent of public instruction, in consultation with the Washington interscholastic activity association, shall promulgate rules by January 1, 1990, regarding loss of eligibility to participate in school-sponsored athletic events for any student athlete found to have violated this chapter. The regents or trustees of each institution of higher education shall promulgate rules by January 1, 1990, regarding loss of eligibility to participate in school-sponsored athletic events for any student athlete found to have violated this chapter. [1989 c 369 § 6.]

69.41.350 Penalties. (1) A person who violates the provisions of this chapter by possessing under two hundred tablets or eight 2cc bottles of steroid without a valid prescription is guilty of a gross misdemeanor.

(2) A person who violates the provisions of this chapter by possessing over two hundred tablets or eight 2cc bottles of steroid without a valid prescription is guilty of a class C felony and shall be punished according to chapter 9A.20 RCW. [2003 c 53 § 326; 1989 c 369 § 4; 1983 1st ex.s. c 4 § 4; 1973 1st ex.s. c 186 § 7. Formerly RCW 69.41.070.]

Intent—Effective date—2003 c 53: See notes following RCW 42.48.180.

Additional notes found at www.leg.wa.gov

69.41.900 Severability—1979 c 110. If any provision of this 1979 act or its application to any person or circumstance is held invalid, the remainder of the act or the applica-
tion of the provision to other persons or circumstances is not affected. [1979 c 110 § 8.]

Chapter 69.43 RCW
PRECURSOR DRUGS

Sections
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69.43.190 Products found at methamphetamine sites—Report.

69.43.010 Report to state board of pharmacy—List of substances—Modification of list—Identification of purchasers—Report of transactions—Penalties. (1) A report to the state board of pharmacy shall be submitted in accordance with this chapter by a manufacturer, wholesaler, retailer, or other person who sells, transfers, or otherwise furnishes to any person any of the following substances or their salts or isomers:
(a) Anthranilic acid;
(b) Barbituric acid;
(c) Chlороphedrine;
(d) Diethyl malonate;
(e) D-lysergic acid;
(f) Ephedrine;
(g) Ergotamine tartrate;
(h) Ethylamine;
(i) Ethyl malonate;
(j) Ethylephedrine;
(k) Lead acetate;
(l) Malonic acid;
(m) Methyleneblue;
(n) Methyleneamid;
(o) Methylphendrine;
(p) Methylpseudoephedrine;
(q) N-acetylanthranilic acid;
(r) Norpseudoephedrine;
(s) Phenylacetic acid;
(t) Phenylpropanolamine;
(u) Piperidine;
(v) Pseudoephedrine; and
(w) Pyrrolidine.
(2) The state board of pharmacy shall administer this chapter and may, by rule adopted pursuant to chapter 34.05 RCW, add a substance to or remove a substance from the list in subsection (1) of this section. In determining whether to add or remove a substance, the board shall consider the following:
(a) The likelihood that the substance is useable as a precursor in the illegal production of a controlled substance as defined in chapter 69.50 RCW;
(b) The availability of the substance;
(c) The relative appropriateness of including the substance in this chapter or in chapter 69.50 RCW; and
(d) The extent and nature of legitimate uses for the substance.
(3)(a) Any manufacturer, wholesaler, retailer, or other person shall, before selling, transferring, or otherwise furnishing any substance specified in subsection (1) of this section to any person, require proper identification from the purchaser.
(b) For the purposes of this subsection, "proper identification" means:
(i) A motor vehicle operator's license or other official state-issued identification of the purchaser containing a photograph of the purchaser, and includes the residential or mailing address of the purchaser, other than a post office box number;
(ii) The motor vehicle license number of any motor vehicle owned or operated by the purchaser;
(iii) A letter of authorization from any business for which any substance specified in subsection (1) of this section is being furnished, which includes the business license number and address of the business;
(iv) A description of how the substance is to be used; and
(v) The signature of the purchaser.
(4) Any manufacturer, wholesaler, retailer, or other person who sells, transfers, or otherwise furnishes the substance specified in subsection (1) of this section to any person shall, not less than twenty-one days before delivery of the substance, submit a report of the transaction, which includes the identification information specified in subsection (3) of this section to the state board of pharmacy. However, the state board of pharmacy may authorize the submission of the reports on a monthly basis with respect to repeated, regular transactions between the furnishers and the recipient involving
the same substance if the state board of pharmacy determines that either of the following exist:

(a) A pattern of regular supply of the substance exists between the manufacturer, wholesaler, retailer, or other person who sells, transfers, or otherwise furnishes such substance and the recipient of the substance; or

(b) The recipient has established a record of using the substance for lawful purposes.

(5) Any person specified in subsection (4) of this section who does not submit a report as required by subsection (4) of this section is guilty of a gross misdemeanor.  [2001 c 96 § 2; 1998 c 245 § 107; 1988 c 147 § 1.]

Intent—2001 c 96: "Communities all over the state of Washington have experienced an increase in the illegal manufacture of methamphetamine. Illegal methamphetamine labs create a significant threat to the health and safety of the people of the state. Some of the chemicals and compounds used to make methamphetamine, and the toxic wastes the process generates, are hazards to the public health. Increases in crime, violence, and the abuse and neglect of children present at laboratory sites are also associated with the increasing number of illegal laboratory sites. The drugs ephedrine, pseudoephedrine, and phenylpropanolamine, which are used in the illegal manufacture of methamphetamine, have been identified as factors in the increase in the number of illegal methamphetamine labs. Therefore, it is the intent of the legislature to place restrictions on the sale and possession of those three drugs in order to reduce the proliferation of illegal methamphetamine laboratories and the associated threats to public health and safety." [2001 c 96 § 1.]

Additional notes found at www.leg.wa.gov

69.43.020 Receipt of substance from source outside state—Report—Penalty.  (1) Any manufacturer, wholesaler, retailer, or other person who receives from a source outside of this state any substance specified in RCW 69.43.010(1) shall submit a report of such transaction to the state board of pharmacy under rules adopted by the board.

(2) Any person specified in subsection (1) of this section who does not submit a report as required by subsection (1) of this section is guilty of a gross misdemeanor.  [2001 c 96 § 3; 1988 c 147 § 2.]

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.030 Exemptions.  RCW 69.43.010 and 69.43.020 do not apply to any of the following:

(1) Any pharmacist or other authorized person who sells or furnishes a substance upon the prescription of a practitioner, as defined in chapter 69.41 RCW;

(2) Any practitioner who administers or furnishes a substance to his or her patients;

(3) Any manufacturer or wholesaler licensed by the state board of pharmacy who sells, transfers, or otherwise furnishes a substance to a licensed pharmacy or practitioner;

(4) Any sale, transfer, furnishing, or receipt of any drug that contains ephedrine, phenylpropanolamine, or pseudoephedrine, or of any cosmetic that contains a substance specified in RCW 69.43.010(1), if such drug or cosmetic is lawfully sold, transferred, or furnished, over the counter without a prescription under chapter 69.04 or 69.41 RCW.  [1988 c 147 § 3.]

69.43.035 Suspicious transactions—Report—Penalty.  (1) Any manufacturer or wholesaler who sells, transfers, or otherwise furnishes any substance specified in RCW 69.43.010(1) to any person in a suspicious transaction shall report the transaction in writing to the state board of pharmacy.

(2) Any person specified in subsection (1) of this section who does not submit a report as required by subsection (1) of this section is guilty of a gross misdemeanor.

(3) For the purposes of this section, "suspicious transaction" means a sale or transfer to which any of the following applies:

(a) The circumstances of the sale or transfer would lead a reasonable person to believe that the substance is likely to be used for the purpose of unlawfully manufacturing a controlled substance under chapter 69.50 RCW, based on such factors as the amount involved, the method of payment, the method of delivery, and any past dealings with any participant in the transaction.  The state board of pharmacy shall adopt by rule criteria for determining whether a transaction is suspicious, taking into consideration the recommendations in appendix A of the report to the United States attorney general by the suspicious orders task force under the federal comprehensive methamphetamine control act of 1996.

(b) The transaction involves payment for any substance specified in RCW 69.43.010(1) in cash or money orders in a total amount of more than two hundred dollars.

(4) The board of pharmacy shall transmit to the department of revenue a copy of each report of a suspicious transaction that it receives under this section.  [2004 c 52 § 6; 2001 c 96 § 4.]

Finding—Severability—Effective date—2004 c 52: See notes following RCW 18.64.044.

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.040 Reporting form.  (1) The department of health, in accordance with rules developed by the state board of pharmacy shall provide a common reporting form for the substances in RCW 69.43.010 that contains at least the following information:

(a) Name of the substance;

(b) Quantity of the substance sold, transferred, or furnished;

(c) The date the substance was sold, transferred, or furnished;

(d) The name and address of the person buying or receiving the substance; and

(e) The name and address of the manufacturer, wholesaler, retailer, or other person selling, transferring, or furnishing the substance.

(2) Monthly reports authorized under RCW 69.43.010(4) may be computer-generated in accordance with rules adopted by the department.  [2001 c 96 § 7; 1989 1st ex.s. c 9 § 441; 1988 c 147 § 4.]

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

Additional notes found at www.leg.wa.gov

69.43.043 Recordkeeping requirements—Penalty.  (1) Any manufacturer or wholesaler who sells, transfers, or otherwise furnishes any substance specified in RCW 69.43.010(1) to any person shall maintain a record of each such sale or transfer.  The records must contain:

(a) The name of the substance;
(b) The quantity of the substance sold, transferred, or furnished;
(c) The date the substance was sold, transferred, or furnished;
(d) The name and address of the person buying or receiving the substance; and
(e) The method of and amount of payment for the substance.

(2) The records of sales and transfers required by this section shall be available for inspection by the state board of pharmacy and its authorized representatives and shall be maintained for two years.

(3) A violation of this section is a gross misdemeanor.

[2001 c 96 § 5.]

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.048 Reporting and recordkeeping requirements—Submission of computer readable data, copies of federal reports. A manufacturer, wholesaler, retailer, or other person who sells, transfers, or otherwise furnishes any substance specified in RCW 69.43.010(1) and who is subject to the reporting or recordkeeping requirements of this chapter may satisfy the requirements by submitting to the state board of pharmacy, and its authorized representatives:

(1) Computer readable data from which all of the required information may be readily derived; or

(2) Copies of reports that are filed under federal law that contain all of the information required by the particular reporting or recordkeeping requirement of this chapter which it is submitted to satisfy. [2001 c 96 § 6.]

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.050 Rules. (1) The state board of pharmacy may adopt all rules necessary to carry out this chapter.

(2) Notwithstanding subsection (1) of this section, the department of health may adopt rules necessary for the administration of this chapter. [1989 1st ex.s. c 9 § 442; 1988 c 147 § 9.]

Additional notes found at www.leg.wa.gov

69.43.060 Theft—Missing quantity—Reporting. (1) The theft or loss of any substance under RCW 69.43.010 discovered by any person regulated by this chapter shall be reported to the state board of pharmacy within seven days after such discovery.

(2) Any difference between the quantity of any substance under RCW 69.43.010 received and the quantity shipped shall be reported to the state board of pharmacy within seven days of the receipt of actual knowledge of the discrepancy. When applicable, any report made pursuant to this subsection shall also include the name of any common carrier or person who transported the substance and the date of shipment of the substance. [1988 c 147 § 6.]

69.43.070 Sale, transfer, or furnishing of substance for unlawful purpose—Receipt of substance with intent to use unlawfully—Class B felony. (1) Any manufacturer, wholesaler, retailer, or other person who sells, transfers, or otherwise furnishes any substance listed in RCW 69.43.010 with knowledge or the intent that the recipient will use the substance unlawfully to manufacture a controlled substance under chapter 69.50 RCW is guilty of a class B felony under chapter 9A.20 RCW.

(2) Any person who receives any substance listed in RCW 69.43.010 with intent to use the substance unlawfully to manufacture a controlled substance under chapter 69.50 RCW is guilty of a class B felony under chapter 9A.20 RCW. [1988 c 147 § 7.]

69.43.080 False statement in report or record—Class C felony. It is unlawful for any person knowingly to make a false statement in connection with any report or record required under this chapter. A violation of this section is a class C felony under chapter 9A.20 RCW. [1988 c 147 § 8.]

69.43.090 Permit to sell, transfer, furnish, or receive substance—Exemptions—Application for permit—Fee—Renewal—Penalty. (1) Any manufacturer, wholesaler, retailer, or other person who sells, transfers, or otherwise furnishes any substance specified in RCW 69.43.010 to any person or who receives from a source outside of the state any substance specified in RCW 69.43.010 shall obtain a permit for the conduct of that business from the state board of pharmacy. However, a permit shall not be required of any manufacturer, wholesaler, retailer, or other person for the sale, transfer, furnishing, or receipt of any drug that contains ephephrine, phenylpropanolamine, or pseudoephedrine, or of any cosmetic that contains a substance specified in RCW 69.43.010(1), if such drug or cosmetic is lawfully sold, transferred, or furnished, or furnished over the counter without a prescription or by a prescription under chapter 69.04 or 69.41 RCW.

(2) Applications for permits shall be filed with the department in writing and signed by the applicant, and shall set forth the name of the applicant, the business in which the applicant is engaged, the business address of the applicant, and a full description of any substance sold, transferred, or otherwise furnished, or received.

(3) The board may grant permits on forms prescribed by it. The permits shall be effective for not more than one year from the date of issuance.

(4) Each applicant shall pay at the time of filing an application for a permit a fee determined by the department.

(5) A permit granted under this chapter may be renewed on a date to be determined by the board, and annually thereafter, upon the filing of a renewal application and the payment of a permit renewal fee determined by the department.

(6) Permit fees charged by the department shall not exceed the costs incurred by the department in administering this chapter.

(7) Selling, transferring, or otherwise furnishing, or receiving any substance specified in RCW 69.43.010 without a required permit, is a gross misdemeanor. [2001 c 96 § 8; 1989 1st ex.s. c 9 § 443; 1988 c 147 § 9.]

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

Additional notes found at www.leg.wa.gov

69.43.100 Refusal, suspension, or revocation of a manufacturer’s or wholesaler’s permit. The board shall
have the power to refuse, suspend, or revoke the permit of any manufacturer or wholesaler upon proof that:

1. The permit was procured through fraud, misrepresentation, or deceit;

2. The permittee has violated or has permitted any employee to violate any of the laws of this state relating to drugs, controlled substances, cosmetics, or nonprescription drugs, or has violated any of the rules and regulations of the board of pharmacy. [1988 c 147 § 10.]

69.43.105 Ephedrine, pseudoephedrine, phenylpropanolamine—Sales restrictions—Record of transaction—Exceptions—Penalty. (1) For purposes of this section, "traditional Chinese herbal practitioner" means a person who is certified as a diplomate in Chinese herbology from the national certification commission for acupuncture and oriental medicine or who has received a certificate in Chinese herbology from a school accredited by the accreditation council on acupuncture and oriental medicine.

(2) A pharmacy licensed by, or shopkeeper or itinerant vendor registered with, the department of health under chapter 18.64 RCW, or an employee thereof, a practitioner as defined in RCW 18.64.011, or a traditional Chinese herbal practitioner may not knowingly sell, transfer, or otherwise furnish to any person a product at retail that he or she knows to contain any detectable quantity of ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers, without first obtaining photo identification of the person that shows the date of birth of the person.

(3) A person buying or receiving a product at retail containing any detectable quantity of ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers, from a pharmacy licensed by, or shopkeeper or itinerant vendor registered with, the department of health under chapter 18.64 RCW, or an employee thereof, a practitioner as defined in RCW 18.64.011, or a traditional Chinese herbal practitioner must first produce photo identification of the person that shows the date of birth of the person.

(4) Any product containing any detectable quantity of ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers, shall be kept (a) behind a counter where the public is not permitted, or (b) in a locked display case so that a customer wanting access must ask an employee of the merchant for assistance.

(5) No pharmacy licensed by, or shopkeeper or itinerant vendor registered with, the department of health under chapter 18.64 RCW, or an employee thereof, a practitioner as defined in RCW 18.64.011, or a traditional Chinese herbal practitioner may sell any product containing any detectable quantity of ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers, to a person that is not at least eighteen years old.

(6) A pharmacy licensed by, or shopkeeper or itinerant vendor registered with, the department of health under chapter 18.64 RCW selling a nonprescription drug containing ephedrine, pseudoephedrine, phenylpropanolamine, or their salts, isomers, or salts of isomers shall require the purchaser to electronically or manually sign a record of the transaction. The record must include the name and address of the purchaser, the date and time of the sale, the name and initials of the shopkeeper, itinerant vendor, pharmacist, pharmacy technician, or employee conducting the transaction, the name of the product being sold, as well as the total quantity in grams, of ephedrine, pseudoephedrine, phenylpropanolamine, or their salts, isomers, or salts of isomers, being sold.

(7) The board of pharmacy, by rule, may exempt products containing ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers, in combination with another active ingredient from the requirements of this section if they are found not to be used in the illegal manufacture of methamphetamine or other controlled dangerous substances. A manufacturer of a drug product may apply for removal of the product from the requirements of this section if the product is determined by the board to have been formulated in such a way as to effectively prevent the conversion of the active ingredient into methamphetamine.

The burden of proof for exemption is upon the person requesting the exemption. The petitioner shall provide the board with evidence that the product has been formulated in such a way as to serve as an effective general deterrent to the conversion of pseudoephedrine into methamphetamine. The evidence must include the furnishing of a valid scientific study, conducted by an independent, professional laboratory and evincing professional quality chemical analysis. Factors to be considered in whether a product should be excluded from this section include but are not limited to:

(a) Ease with which the product can be converted to methamphetamine;

(b) Ease with which ephedrine, pseudoephedrine, or phenylpropanolamine is extracted from the substance and whether it forms an emulsion, salt, or other form;

(c) Whether the product contains a "molecular lock" that renders it incapable of being converted into methamphetamine;

(d) Presence of other ingredients that render the product less likely to be used in the manufacture of methamphetamine;

(e) Any pertinent data that can be used to determine the risk of the substance being used in the illegal manufacture of methamphetamine or any other controlled substance.

(8) Nothing in this section applies:

(a) To any product containing ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers that is not the only active ingredient and that is in liquid, liquid capsule, or gel capsule form;

(b) To the sale of a product that may only be sold upon the presentation of a prescription;

(c) To the sale of a product by a traditional Chinese herbal practitioner to a patient; or

(d) When the details of the transaction are recorded in a pharmacy profile individually identified with the recipient and maintained by a licensed pharmacy.

(9)(a) No pharmacy licensed by, or shopkeeper or itinerant vendor registered with, the department of health under chapter 18.64 RCW, a practitioner as defined in RCW 18.64.011, or a traditional Chinese herbal practitioner may retaliate against any employee that has made a good faith attempt to comply with the requirements of this section by requesting that a customer present photo identification, making a reasonable effort to determine the customer’s age.

(b) No pharmacy licensed by, or shopkeeper or itinerant vendor registered with, the department of health under chap-
ter 18.64 RCW, a practitioner as defined in RCW 18.64.011, or a traditional Chinese herbal practitioner is subject to prosecution under subsection (10) of this section if they made a good faith attempt to comply with the requirements of this section by requesting that a customer present photo identification, making a reasonable effort to determine the customer’s age.

(10) A violation of this section is a gross misdemeanor. [2010 c 182 § 1; 2005 c 388 § 2.]

Finding—2005 c 388: "Restricting access to certain precursor drugs used to manufacture methamphetamine to ensure that they are only sold at retail to individuals who will use them for legitimate purposes upon production of proper identification is an essential step to controlling the manufacture of methamphetamine." [2005 c 388 § 1.]

Effective dates—2005 c 388: "(1) Section 2 of this act takes effect October 1, 2005.
(2) Sections 1, 3 through 7, 9, and 10 of this act take effect January 1, 2006.
(3) Section 8 of this act is necessary for the immediate preservation of public peace, health, or safety, or support of the state government and its existing public institutions, and takes effect immediately [May 11, 2005]." [2005 c 388 § 11.]

Severability—2005 c 388: "If any provision of this act or its application to any person or circumstance is held invalid, the remainder of the act or the application of the provision to other persons or circumstances is not affected." [2005 c 388 § 10.]

69.43.110 Ephedrine, pseudoephedrine, phenylpropanolamine—Sales restrictions—Electronic sales tracking system—Penalty. (1) It is unlawful for a pharmacy licensed by, or shopkeeper or itinerant vendor registered with, the department of health under chapter 18.64 RCW, or an employee thereof, or a practitioner as defined in RCW 18.64.011, knowingly to sell, transfer, or to otherwise furnish, in a single transaction a total of more than 3.6 grams of ephedrine, pseudoephedrine, or phenylpropanolamine, their salts, isomers, or salts of isomers, in any twenty-four hour period or more than a total of nine grams per purchaser in any thirty-day period.

(2) It is unlawful for a person who is not a manufacturer, wholesaler, pharmacy, practitioner, shopkeeper, or itinerant vendor licensed by or registered with the department of health under chapter 18.64 RCW to purchase or acquire more than 3.6 grams in any twenty-four hour period, or more than a total of nine grams in any thirty-day period, of the substances specified in subsection (1) of this section.

(3) It is unlawful for any person to sell or distribute any of the substances specified in subsection (1) of this section unless the person is licensed by or registered with the department of health under chapter 18.64 RCW, or is a practitioner as defined in RCW 18.64.011.

(4)(a) Beginning July 1, 2011, or the date upon which the electronic sales tracking system established under RCW 69.43.165 is available, whichever is later, a pharmacy licensed by, or shopkeeper or itinerant vendor registered with, the department of health under chapter 18.64 RCW shall, before completing a sale under this section, submit the required information to the electronic sales tracking system established under RCW 69.43.165, as long as such a system is available without cost to the pharmacy, shopkeeper, or itinerant vendor for accessing the system. The pharmacy, shopkeeper, or itinerant vendor may not complete the sale if the system generates a stop sale alert, except as permitted in RCW 69.43.165.

(b) If a pharmacy, shopkeeper, or itinerant vendor selling a nonprescription drug containing ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers experiences mechanical or electronic failure of the electronic sales tracking system and is unable to comply with the electronic sales tracking requirement, he or she shall maintain a written log or an alternative electronic record-keeping mechanism until such time as he or she is able to comply with the electronic sales tracking requirement.

(c) A pharmacy, shopkeeper, or itinerant vendor selling a nonprescription drug containing ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers may seek an exemption from submitting transactions to the electronic sales tracking system in writing to the board of pharmacy stating the reasons for the exemption. The board may grant an exemption for good cause shown, but in no event shall a granted exemption exceed one hundred eighty days. The board may grant multiple exemptions for any pharmacy, shopkeeper, or itinerant vendor if the good cause shown indicates significant hardship for compliance with this section. A pharmacy, shopkeeper, or itinerant vendor that receives an exemption shall maintain a logbook in hardcopy form and must require the purchaser to provide the information required under this section before the completion of any sale. The logbook shall be maintained as a record of each sale for inspection by any law enforcement officer or board inspector during normal business hours in accordance with any rules adopted pursuant to RCW 69.43.165. For purposes of this subsection (4)(c), "good cause" includes, but is not limited to, situations where the installation of the necessary equipment to access the system is unavailable or cost prohibitive to the pharmacy, shopkeeper, or itinerant vendor.

(d) A pharmacy, shopkeeper, or itinerant vendor may withdraw from participating in the electronic sales tracking system if the system is no longer being furnished without cost for accessing the system. A pharmacy, shopkeeper, or itinerant vendor who withdraws from the electronic sales tracking system is subject to the same requirements as a pharmacy, shopkeeper, or itinerant vendor who has been granted an exemption under (c) of this subsection.

(e) For the purposes of this subsection (4) and RCW 69.43.165:

(i) "Cost for accessing the system" means costs relating to:
(A) Access to the web-based electronic sales tracking software, including inputting and retrieving data;
(B) The web-based software known as software as a service;
(C) Training; and
(D) Technical support to integrate to point of sale vendors, if necessary.

(ii) "Cost for accessing the system" does not include:
(A) Costs relating to required internet access;
(B) Optional hardware that a pharmacy may choose to purchase for work flow purposes; or
(C) Other equipment.

(5) A violation of this section is a gross misdemeanor. [2010 c 182 § 2; 2005 c 388 § 4; 2004 c 52 § 5; 2001 c 96 § 9.]
Finding—Effective dates—Severability—2005 c 388: See notes following RCW 69.43.105.

Finding—Severability—Effective date—2004 c 52: See notes following RCW 18.64.044.

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.120 Ephedrine, pseudoephedrine, phenylpropanolamine—Possession of more than fifteen grams—Penalty—Exceptions. (1) Any person who possesses more than fifteen grams of ephedrine, pseudoephedrine, or phenylpropanolamine, their salts, isomers, or salts of isomers, or a combination of any of those substances, is guilty of a gross misdemeanor.

(2) This section does not apply to any of the following:

(a) A pharmacist or other authorized person who sells or furnishes ephedrine, pseudoephedrine, or phenylpropanolamine, their salts, isomers, or salts of isomers upon the prescription of a practitioner, as defined in RCW 69.41.010;

(b) A practitioner who administers or furnishes ephedrine, pseudoephedrine, or phenylpropanolamine, their salts, isomers, or salts of isomers to his or her patients;

(c) A pharmacy, manufacturer, or wholesaler licensed by, or shopkeeper or itinerant vendor registered with, the department of health under chapter 18.64 RCW;

(d) A person in the course of his or her business of selling, transporting, or storing ephedrine, pseudoephedrine, or phenylpropanolamine, their salts, isomers, or salts of isomers, for a person described in (a), (b), or (c) of this subsection; or

(e) A person in possession of more than fifteen grams of ephedrine, pseudoephedrine, or phenylpropanolamine, their salts, isomers, or salts of isomers in their home or residence under circumstances consistent with typical medicinal or household use as indicated by, but not limited to, storage location and possession of products in a variety of strengths, brands, types, purposes, and expiration dates. [2001 c 96 § 10.]

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.130 Exemptions—Pediatric products—Products exempted by the state board of pharmacy. RCW 69.43.110 and 69.43.120 do not apply to:

(1) Pediatric products primarily intended for administration to children under twelve years of age, according to label instructions, either: (a) In solid dosage form whose individual dosage units do not exceed fifteen milligrams of ephedrine, pseudoephedrine, or phenylpropanolamine; or (b) in liquid form whose recommended dosage, according to label instructions, does not exceed fifteen milligrams of ephedrine, pseudoephedrine, or phenylpropanolamine per five milliliters of liquid product;

(2) Pediatric liquid products primarily intended for administration to children under two years of age for which the recommended dosage does not exceed two milliliters and the total package content does not exceed one fluid ounce;

(3) Products that the state board of pharmacy, upon application of a manufacturer, exempts by rule from RCW 69.43.110 and 69.43.120 because the product has been formulated in such a way as to effectively prevent the conversion of the active ingredient into methamphetamine, or its salts or precursors; or

(4) Products, as packaged, that the board of pharmacy, upon application of a manufacturer, exempts from RCW *69.43.110(1)(b) and 69.43.120 because:

(a) The product meets the federal definition of an ordinary over-the-counter pseudoephedrine product as defined in 21 U.S.C. 802;

(b) The product is a salt, isomer, or salts of isomers of pseudoephedrine and, as packaged, has a total weight of more than three grams but the net weight of the pseudoephedrine base is equal to or less than three grams; and

(c) The board of pharmacy determines that the value to the people of the state of having the product, as packaged, available for sale to consumers outweighs the danger, and the product, as packaged, has not been used in the illegal manufacture of methamphetamine. [2004 c 52 § 7; 2001 c 96 § 11.]

*Reviser’s note: RCW 69.43.110 was amended by 2010 c 182 § 2, changing subsection (1)(b) to subsection (1).

Finding—Severability—Effective date—2004 c 52: See notes following RCW 18.64.044.

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.135 Iodine, methylsulfonylmethane—Sales restrictions—Recording of transactions—Penalties. (1) The definitions in this subsection apply throughout this section unless the context clearly requires otherwise.

(a) "Iodine matrix" means iodine at a concentration greater than two percent by weight in a matrix or solution.

(b) "Matrix" means something, as a substance, in which something else originates, develops, or is contained.

(c) "Methylsulfonylmethane" means methylsulfonylmethane in its powder form only, and does not include products containing methylsulfonylmethane in other forms such as liquids, tablets, capsules not containing methylsulfonylmethane in pure powder form, ointments, creams, cosmetics, foods, and beverages.

(2) Any person who knowingly purchases in a thirty-day period or possesses any quantity of iodine in its elemental form, an iodine matrix, or more than two pounds of methylsulfonylmethane is guilty of a gross misdemeanor, except as provided in subsection (3) of this section.

(3) Subsection (2) of this section does not apply to:

(a) A person who possesses iodine in its elemental form or an iodine matrix as a prescription drug, under a prescription issued by a licensed veterinarian, physician, or advanced registered nurse practitioner;

(b) A person who possesses iodine in its elemental form, an iodine matrix, or any quantity of methylsulfonylmethane in its powder form and is actively engaged in the practice of animal husbandry of livestock;

(c) A person who possesses iodine in its elemental form or an iodine matrix in conjunction with experiments conducted in a chemistry or chemistry-related laboratory maintained by a:

(i) Public or private secondary school;

(ii) Public or private institution of higher education that is accredited by a regional or national accrediting agency recognized by the United States department of education;
(iii) Manufacturing facility, government agency, or research facility in the course of lawful business activities;

(d) A veterinarian, physician, advanced registered nurse practitioner, pharmacist, retailer, distributor, wholesaler, manufacturer, warehouse operator, or common carrier, or an agent of any of these persons who possesses iodine in its elemental form, an iodine matrix, or methylsulfonylmethane in its powder form in the regular course of lawful business activities; or

(e) A person working in a general hospital who possesses iodine in its elemental form or an iodine matrix in the regular course of employment at the hospital.

(4) Any person who purchases any quantity of iodine in its elemental form, an iodine matrix, or any quantity of methylsulfonylmethane must present an identification card or driver’s license issued by any state in the United States or jurisdiction of another country before purchasing the item.

(5) The Washington state patrol shall develop a form to be used in recording transactions involving iodine in its elemental form, an iodine matrix, or methylsulfonylmethane. A person who sells or otherwise transfers any quantity of iodine in its elemental form, an iodine matrix, or any quantity of methylsulfonylmethane to a person for any purpose authorized in subsection (3) of this section must record each sale or transfer. The record must be made on the form developed by the Washington state patrol and must be retained by the person for at least three years. The Washington state patrol or any local law enforcement agency may request access to the records.

(a) Failure to make or retain a record required under this subsection is a misdemeanor.

(b) Failure to comply with a request for access to records required under this subsection to the Washington state patrol or a local law enforcement agency is a misdemeanor. [2011 c 336 § 838; 2006 c 188 § 1.]

69.43.140 Civil penalty—State board of pharmacy waiver. (1) In addition to the other penalties provided for in this chapter or in chapter 18.64 RCW, the state board of pharmacy may impose a civil penalty, not to exceed ten thousand dollars for each violation, on any licensee or registrant who has failed to comply with this chapter or the rules adopted under this chapter. In the case of a continuing violation, every day the violation continues shall be considered a separate violation.

(2) The state board of pharmacy may waive the suspension or revocation of a license or registration issued under chapter 18.64 RCW, or waive any civil penalty under this chapter, if the licensee or registrant establishes that he or she acted in good faith to prevent violations of this chapter, and the violation occurred despite the licensee’s or registrant’s exercise of due diligence. In making such a determination, the state board of pharmacy may consider evidence that an employer trained employees on how to sell, transfer, or otherwise furnish substances specified in RCW 69.43.010(1) in accordance with applicable laws. [2001 c 96 § 12.]

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.150 Application of chapter to local government. This chapter is applicable and uniform throughout this state and in all counties, cities, code cities, and towns therein. A county, city, code city, or town may not adopt or enforce any ordinance, pertaining to this chapter, which prohibits conduct that is not prohibited under this chapter, or defining violations or penalties different from those provided under this chapter. However, this section does not preclude a county, city, code city, or town from revoking, canceling, suspending, or otherwise limiting a business or professional license if it is issued for conduct that violates any provision of this chapter. [2001 c 96 § 13.]

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.160 Ephedrine, pseudoephedrine, phenylpropanolamine—Methods to prevent sales violations—Department of health preparation of sign summarizing prohibitions. (1) To prevent violations of RCW 69.43.110, every licensee and registrant under chapter 18.64 RCW, who sells at retail any products containing ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers, shall do either or may do both of the following:

(a) Program scanners, cash registers, or other electronic devices used to record sales in a manner that will alert persons handling transactions to potential violations of RCW 69.43.110(1) and/or prevent such violations; or

(b) Place one or more signs on the premises to notify customers of the prohibitions of RCW 69.43.110. Any such sign may, but is not required to, conform to the language and format prepared by the department of health under subsection (2) of this section.

(2) The department of health shall prepare language and format for a sign summarizing the prohibitions in RCW 69.43.110 and 69.43.120 and make the language and format available to licensees and registrants under chapter 18.64 RCW, for voluntary use in their places of business to inform customers and employees of the prohibitions. Nothing in this section requires the department of health to provide licensees or registrants with copies of signs, or any licensee or registrant to use the specific language or format prepared by the department under this subsection. [2001 c 96 § 14.]

Intent—Severability—2001 c 96: See notes following RCW 69.43.010.

69.43.165 Ephedrine, pseudoephedrine, phenylpropanolamine—Electronic sales tracking system—Board of pharmacy authority to adopt rules. (1) The board of pharmacy shall implement a real-time electronic sales tracking system to monitor the nonprescription sale of products in this state containing any detectable quantity of ephedrine, pseudoephedrine, phenylpropanolamine, or their salts, isomers, or salts of isomers, provided that the system is available to the state without cost for accessing the system to the state or retailers. The board is authorized to enter into a public-private partnership, through a memorandum of understanding or similar arrangement, to make the system available.

(2) The records submitted to the tracking system are for the confidential use of the pharmacy, shopkeeper, or itinerant vendor who submitted them, except that:

(a) The records must be produced in court when lawfully required;
(b) The records must be open for inspection by the board of pharmacy; and

c) The records must be available to any general or limited authority Washington peace officer to enforce the provisions of this chapter or to federal law enforcement officers in accordance with rules adopted by the board of pharmacy regarding the privacy of the purchaser of products covered by chapter 182, Laws of 2010 and law enforcement access to the records submitted to the tracking system as provided in this section consistent with the federal combat meth act.

(3) The electronic sales tracking system shall be capable of generating a stop sale alert, which shall be a notification that completion of the sale would result in the seller or purchaser violating the quantity limits in RCW 69.43.110 (1) and (2). The system shall contain an override function for use by a dispenser of ephedrine, pseudoephedrine, phenylpropanolamine, or their salts, isomers, or salts of isomers, who has a reasonable fear of imminent bodily harm. Each instance in which the override function is utilized shall be logged by the system.

(4) The board of pharmacy shall have the authority to adopt rules necessary to implement and enforce the provisions of this section. The board of pharmacy shall adopt rules regarding the privacy of the purchaser of products covered by chapter 182, Laws of 2010, and any public or law enforcement access to the records submitted to the tracking system as provided in subsection (2)(c) of this section consistent with the federal combat meth act.

(5) The board of pharmacy may not raise licensing or registration fees to fund the rule making or implementation of this section. [2010 c 182 § 3.]

69.43.180 Expansion of log requirements—Petition by law enforcement. (1) The Washington association of sheriffs and police chiefs or the Washington state patrol may petition the state board of pharmacy to apply the log requirements in *RCW 69.43.170 to one or more products that contain ephedrine, pseudoephedrine, or phenylpropanolamine, or their salts, isomers, or salts of isomers, that is not the only active ingredient and that is in liquid, liquid capsule, or gel capsule form. The petition shall establish that:

(a) Ephedrine, pseudoephedrine, or phenylpropanolamine can be effectively extracted from the product and converted into methamphetamine or another controlled dangerous substance; and

(b) Law enforcement, the Washington state patrol, or the department of ecology are finding substantial evidence that the product is being used for the illegal manufacture of methamphetamine or another controlled dangerous substance.

(2) The board of pharmacy shall adopt rules when a petition establishes that requiring the application of the log requirements in *RCW 69.43.170 to the sale of the product at retail is warranted based upon the effectiveness and extent of use of the product for the illegal manufacture of methamphetamine or other controlled dangerous substances and the extent of the burden of any restrictions upon consumers. The board of pharmacy may adopt emergency rules to apply the log requirements to the sale of a product when the petition establishes that the immediate restriction of the product is necessary in order to protect public health and safety. [2005 c 388 § 3.]

Finding—Effective dates—Severability—2005 c 388: See notes following RCW 69.43.105.

69.43.190 Products found at methamphetamine sites—Report. Each county sheriff shall compile and maintain a record of commercial products containing ephedrine, pseudoephedrine, or phenylpropanolamine and packaging found at methamphetamine laboratory sites. The data shall be forwarded to the Washington association of sheriffs and police chiefs and shall be reported to the legislature by November 1, 2007, and annually thereafter. [2005 c 388 § 9.]

Finding—Effective dates—Severability—2005 c 388: See notes following RCW 69.43.105.

Chapter 69.45 RCW

DRUG SAMPLES

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69.45.050 Distribution of drug samples—Written request—No fee or charge permitted—Possession of legend drugs or controlled substances by manufacturers’ representatives.

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69.45.070 Registration fees—Penalty.

69.45.080 Violations of chapter—Manufacturer’s liability—Penalty—Seizure of drug samples.

69.45.090 Confidentiality.

69.45.900 Severability—1987 c 411.

69.45.010 Definitions. The definitions in this section apply throughout this chapter.

(1) "Board" means the board of pharmacy.

(2) "Drug samples" means any federal food and drug administration approved controlled substance, legend drug, or products requiring prescriptions in this state, which is distributed at no charge to a practitioner by a manufacturer or a manufacturer’s representative, exclusive of drugs under clinical investigations approved by the federal food and drug administration.

(3) "Controlled substance" means a drug, substance, or immediate precursor of such drug or substance, so designated
under or pursuant to chapter 69.50 RCW, the uniform controlled substances act.

(4) "Deliver" or "delivery" means the actual, constructive, or attempted transfer from one person to another of a drug or device, whether or not there is an agency relationship.

(5) "Dispense" means the interpretation of a prescription or order for a drug, biological, or device and, pursuant to that prescription or order, the proper selection, measuring, compounding, labeling, or packaging necessary to prepare that prescription or order for delivery.

(6) "Distribute" means to deliver, other than by administering or dispensing, a legend drug.

(7) "Legend drug" means any drug that is required by state law or by regulations of the board to be dispensed on prescription only or is restricted to use by practitioners only.

(8) "Manufacturer" means a person or other entity engaged in the manufacture or distribution of drugs or devices, but does not include a manufacturer’s representative.

(9) "Person" means any individual, corporation, governmental or governmental subdivision or agency, business trust, estate, trust, partnership, association, or any other legal entity.

(10) "Practitioner" means a physician under chapter 18.71 RCW, an osteopathic physician or an osteopathic physician and surgeon under chapter 18.57 RCW, a dentist under chapter 18.32 RCW, a podiatric physician and surgeon under chapter 18.22 RCW, a veterinarian under chapter 18.92 RCW, a pharmacist under chapter 18.64 RCW, a commissioned medical or dental officer in the United States armed forces or the public health service in the discharge of his or her official duties, a duly licensed physician or dentist employed by the veterans administration in the discharge of his or her official duties, a registered nurse or advanced registered nurse practitioner under chapter 18.79 RCW when authorized to prescribe by the nursing care quality assurance commission, an osteopathic physician assistant under chapter 18.57A RCW when authorized by the board of osteopathic medicine and surgery, or a physician assistant under chapter 18.71A RCW when authorized by the medical quality assurance commission.

(11) "Manufacturer’s representative" means an agent or employee of a drug manufacturer who is authorized by the drug manufacturer to possess drug samples for the purpose of distribution in this state to appropriately authorized health care practitioners.

(12) "Reasonable cause" means a state of facts found to exist that would warrant a reasonably intelligent and prudent person to believe that a person has violated state or federal drug laws or regulations.

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

(14) "Secretary" means the secretary of health or the secretary’s designee. [1994 sp.s. c 9 § 738; 1989 1st ex.s. c 9 § 444; 1987 c 411 § 1.]
expiration dates. (1) Drug samples shall be stored in compliance with the requirements of federal and state laws, rules, and regulations.

(2) Drug samples shall be maintained in a locked area to which access is limited to persons authorized by the manufacturer.

(3) Drug samples shall be stored and transported in such a manner as to be free of contamination, deterioration, and adulteration.

(4) Drug samples shall be stored under conditions of temperature, light, moisture, and ventilation so as to meet the label instructions for each drug.

(5) Drug samples which have exceeded the expiration date shall be physically separated from other drug samples until disposed of or returned to the manufacturer. [1987 c 411 § 4.]

69.45.050 Distribution of drug samples—Written request—No fee or charge permitted—Possession of legend drugs or controlled substances by manufacturers’ representatives. (1) Drug samples may be distributed by a manufacturer or a manufacturer’s representative only to practitioners legally authorized to prescribe such drugs or, at the request of such practitioner, to pharmacies of hospitals or other health care entities. The recipient of the drug sample must execute a written receipt upon delivery that is returned to the manufacturer or the manufacturer’s representative.

(2) Drug samples may be distributed by a manufacturer or a manufacturer’s representative only to a practitioner legally authorized to prescribe such drugs pursuant to a written request for such samples. The request shall contain:

(a) The recipient’s name, address, and professional designation;

(b) The name, strength, and quantity of the drug samples delivered;

(c) The name or identification of the manufacturer and of the individual distributing the drug sample; and

(d) The dated signature of the practitioner requesting the drug sample.

(3) No fee or charge may be imposed for sample drugs distributed in this state.

(4) A manufacturer’s representative shall not possess legend drugs or controlled substances other than those distributed by the manufacturer they represent. Nothing in this section prevents a manufacturer’s representative from possessing a legally prescribed and dispensed legend drug or controlled substance. [1989 c 164 § 2; 1987 c 411 § 5.]

Legislative finding—1989 c 164: “The legislature finds that chapter 69.45 RCW is more restrictive than the federal prescription drug marketing act of 1987, and the legislature further finds that a change in chapter 69.45 RCW accepting the position of the federal law is beneficial to the citizens of this state.” [1989 c 164 § 1.]

69.45.060 Disposal of surplus, outdated, or damaged drug samples. Surplus, outdated, or damaged drug samples shall be disposed of as follows:

(1) Returned to the manufacturer; or

(2) Witnessed destruction by such means as to assure that the drug cannot be retrieved. However, controlled substances shall be returned to the manufacturer or disposed of in accordance with rules adopted by the board: PROVIDED, That the board shall adopt by rule the regulations of the federal drug enforcement administration or its lawful successor unless, stating reasonable grounds, it adopts rules consistent with such regulations. [1987 c 411 § 6.]

69.45.070 Registration fees—Penalty. The department may charge reasonable fees for registration. The registration fee shall not exceed the fee charged by the department for a pharmacy location license. If the registration fee is not paid on or before the date due, a renewal or new registration may be issued only upon payment of the registration renewal fee and a penalty fee equal to the registration renewal fee. [1991 c 229 § 8; 1989 1st ex.s. c 9 § 447; 1987 c 411 § 7.]

Additional notes found at www.leg.wa.gov

69.45.080 Violations of chapter—Manufacturer’s liability—Penalty—Seizure of drug samples. (1) The manufacturer is responsible for the actions and conduct of its representatives with regard to drug samples.

(2) The board may hold a public hearing to examine a possible violation and may require a designated representative of the manufacturer to attend.

(3) If a manufacturer fails to comply with this chapter following notification by the board, the board may impose a civil penalty of up to five thousand dollars. The board shall take no action to impose any civil penalty except pursuant to a hearing held in accordance with chapter 34.05 RCW.

(4) Specific drug samples which are distributed in this state in violation of this chapter, following notification by the board, shall be subject to seizure following the procedures set out in RCW 69.41.060. [1987 c 411 § 8.]

69.45.090 Confidentiality. All records, reports, and information obtained by the board from or on behalf of a manufacturer or manufacturer’s representative under this chapter are confidential and exempt from public inspection and copying under chapter 42.56 RCW. This section does not apply to public disclosure of the identity of persons found by the board to have violated state or federal law, rules, or regulations. This section is not intended to restrict the investigations and proceedings of the board so long as the board maintains the confidentiality required by this section. [2005 c 274 § 330; 1987 c 411 § 9.]

Part headings not law—Effective date—2005 c 274: See RCW 42.56.901 and 42.56.902.

69.45.900 Severability—1987 c 411. If any provision of this act or its application to any person or circumstance is held invalid, the remainder of the act or the application of the provision to other persons or circumstances is not affected. [1987 c 411 § 12.]

Chapter 69.50 RCW

UNIFORM CONTROLLED SUBSTANCES ACT

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(iv) any substance to the extent not intended for human consumption before an exemption takes effect with respect to the substance.

(f) "Deliver" or "delivery," means the actual or constructive transfer from one person to another of a substance, whether or not there is an agency relationship.

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

(h) "Dispense" means the interpretation of a prescription or order for a controlled substance and, pursuant to that prescription or order, the proper selection, measuring, compounding, labeling, or packaging necessary to prepare that prescription or order for delivery.

(i) "Dispenser" means a practitioner who dispenses.

(j) "Distribute" means to deliver other than by administering or dispensing a controlled substance.

(k) "Distributor" means a person who distributes.

(l) "Drug" means (1) a controlled substance recognized as a drug in the official United States pharmacopoeia/national formulary or the official homeopathic pharmacopoeia of the United States, or any supplement to them; (2) controlled substances intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in individuals or animals; (3) controlled substances (other than food) intended to affect the structure or any function of the body of individuals or animals; and (4) controlled substances intended for use as a component of any article specified in (1), (2), or (3) of this subsection. The term does not include devices or their components, parts, or accessories.

(m) "Drug enforcement administration" means the drug enforcement administration in the United States Department of Justice, or its successor agency.

(n) "Electronic communication of prescription information" means the communication of prescription information by computer, or the transmission of an exact visual image of a prescription by facsimile, or other electronic means for original prescription information or prescription refill information for a Schedule III-V controlled substance between an authorized practitioner and a pharmacy or the transfer of prescription information for a controlled substance from one pharmacy to another pharmacy.

(o) "Immediate precursor" means a substance:

(1) that the state board of pharmacy has found to be and by rule designates as being the principal compound commonly used, or produced primarily for use, in the manufacture of a controlled substance; and

(2) that is an immediate chemical intermediary used or likely to be used in the manufacture of a controlled substance; and

(3) the control of which is necessary to prevent, curtail, or limit the manufacture of the controlled substance.

(p) "Isomer" means an optical isomer, but in RCW 69.50.101 (r)(5), 69.50.204(a) (12) and (34), and 69.50.206(b)(4), the term includes any geometrical isomer; in RCW 69.50.204(a) (8) and (42), and 69.50.210(c) the term includes any positional isomer; and in RCW 69.50.204(a)(35), 69.50.204(c), and 69.50.208(a) the term includes any positional or geometric isomer.

(q) "Manufacture" means the production, preparation, propagation, compounding, conversion, or processing of a controlled substance, either directly or indirectly or by extraction from substances of natural origin, or independently by means of chemical synthesis, or by a combination of extraction and chemical synthesis, and includes any packaging or repackaging of the substance or labeling or relabeling of its container. The term does not include the preparation, compounding, packaging, repackaging, labeling, or relabeling of a controlled substance:

(1) by a practitioner as an incident to the practitioner’s administering or dispensing of a controlled substance in the course of the practitioner’s professional practice; or

(2) by a practitioner, or by the practitioner’s authorized agent under the practitioner’s supervision, for the purpose of, or as an incident to, research, teaching, or chemical analysis and not for sale.

(r) "Marijuana" or "marihuana" means all parts of the plant Cannabis, whether growing or not; the seeds thereof; the resin extracted from any part of the plant; and every compound, manufacture, salt, derivative, mixture, or preparation of the plant, its seeds or resin. The term does not include the mature stalks of the plant, fiber produced from the stalks, oil or cake made from the seeds of the plant, any other compound, manufacture, salt, derivative, mixture, or preparation of the mature stalks (except the resin extracted therefrom), fiber, oil, or cake, or the sterilized seed of the plant which is incapable of germination.

(s) "Narcotic drug" means any of the following, whether produced directly or indirectly by extraction from substances of vegetable origin, or independently by means of chemical synthesis, or by a combination of extraction and chemical synthesis:

(1) Opium, opium derivative, and any derivative of opium or opium derivative, including their salts, isomers, and salts of isomers, whenever the existence of the salts, isomers, and salts of isomers is possible within the specific chemical designation. The term does not include the isoquinoline alkaloids of opium.

(2) Synthetic opiate and any derivative of synthetic opiate, including their isomers, esters, ethers, salts, and salts of isomers, esters, ethers, and salts when the existence of the isomers, esters, ethers, and salts is possible within the specific chemical designation.

(3) Poppy straw and concentrate of poppy straw.

(4) Coca leaves, except coca leaves and extracts of coca leaves from which cocaine, ecolgonine, and derivatives or ecolgonine or their salts have been removed.

(5) Cocaine, or any salt, isomer, or salt of isomer thereof.

(6) Cocaine base.

(7) Ecolgonine, or any derivative, salt, isomer, or salt of isomer thereof.

(8) Any compound, mixture, or preparation containing any quantity of any substance referred to in subparagraphs (1) through (7).

(t) "Opiate" means any substance having an addiction-forming or addiction-sustaining liability similar to morphine or being capable of conversion into a drug having addiction-forming or addiction-sustaining liability. The term includes opium, substances derived from opium (opium derivatives), and synthetic opiates. The term does not include, unless specifically designated as controlled under RCW 69.50.201, the dextrorotatory isomer of 3-methoxy-n-methylmorphinan and its salts (dextromethorphan). The term includes the racemic and levorotatory forms of dextromethorphan.
(u) "Opium poppy" means the plant of the species Papaver somniferum L., except its seeds.

(v) "Person" means individual, corporation, business trust, estate, trust, partnership, association, joint venture, government, governmental subdivision or agency, or any other legal or commercial entity.

(w) "Poppy straw" means all parts, except the seeds, of the opium poppy, after mowing.

(x) "Practitioner" means:

(1) A physician under chapter 18.71 RCW; a physician assistant under chapter 18.71A RCW; an osteopathic physician and surgeon under chapter 18.57 RCW; an osteopathic physician assistant under chapter 18.57A RCW who is licensed under RCW 18.57A.020 subject to any limitations in RCW 18.57A.040; an optometrist licensed under chapter 18.53 RCW who is certified by the optometry board under RCW 18.53.010 subject to any limitations in RCW 18.53.010; a dentist under chapter 18.32 RCW; a podiatric physician and surgeon under chapter 18.22 RCW; a veterinarian under chapter 18.92 RCW; a registered nurse, advanced registered nurse practitioner, orlicensed practical nurse under chapter 18.79 RCW; a naturopathic physician under chapter 18.36A RCW who is licensed under RCW 18.36A.030 subject to any limitations in RCW 18.36A.040; a pharmacist under chapter 18.64 RCW or a scientific investigator under this chapter, licensed, registered or otherwise permitted insofar as is consistent with those licensing laws to distribute, dispense, conduct research with respect to or administer a controlled substance in the course of their professional practice or research in this state.

(2) A pharmacy, hospital or other institution licensed, registered, or otherwise permitted to distribute, dispense, conduct research with respect to or to administer a controlled substance in the course of professional practice or research in this state.

(3) A physician licensed to practice medicine and surgery, a physician licensed to practice osteopathic medicine and surgery, a dentist licensed to practice dentistry, a podiatric physician and surgeon licensed to practice podiatric medicine and surgery, an advanced registered nurse practitioner licensed to prescribe controlled substances, or a veterinarian licensed to practice veterinary medicine in any state of the United States.

(y) "Prescription" means an order for controlled substances issued by a practitioner duly authorized by law or rule in the state of Washington to prescribe controlled substances within the scope of his or her professional practice for a legitimate medical purpose.

(z) "Production" includes the manufacturing, planting, cultivating, growing, or harvesting of a controlled substance.

(aa) "Secretary" means the secretary of health or the secretary's designee.

(bb) "State," unless the context otherwise requires, means a state of the United States, the District of Columbia, the Commonwealth of Puerto Rico, or a territory or insular possession subject to the jurisdiction of the United States.

(cc) "Ultimate user" means an individual who lawfully possesses a controlled substance for the individual’s own use or for the use of a member of the individual’s household or for administering to an animal owned by the individual or by a member of the individual’s household. [2012 c 8 § 1; 2010 c 177 § 1; 2003 c 142 § 4; 1998 c 222 § 3; 1996 c 178 § 18; 1994 sp.s. c 9 § 739; 1993 c 187 § 1. Prior: 1990 c 248 § 1; 1990 c 219 § 3; 1990 c 196 § 8; 1989 1st ex.s. c 9 § 429; 1987 c 144 § 2; 1986 c 124 § 1; 1984 c 153 § 18; 1980 c 71 § 2; 1973 2nd ex.s. c 38 § 1; 1971 ex.s. c 308 § 69.50.101.]

Reviser's note: The definitions in this section have been alphabetized pursuant to RCW 1.08.015(2)(k).

Severability—2003 c 142: See note following RCW 18.53.010.

Finding—1990 c 219: See note following RCW 69.41.030.

Additional notes found at www.leg.wa.gov

69.50.102 Drug paraphernalia—Definitions. (a) As used in this chapter, "drug paraphernalia" means all equipment, products, and materials of any kind which are used, intended for use, or designed for use in planting, propagating, cultivating, growing, harvesting, manufacturing, compounding, converting, producing, processing, preparing, testing, analyzing, packaging, repackaging, storing, containing, concealing, injecting, ingesting, inhaling, or otherwise introducing into the human body a controlled substance. It includes, but is not limited to:

(1) Kits used, intended for use, or designed for use in planting, propagating, cultivating, growing, or harvesting of any species of plant which is a controlled substance or from which a controlled substance can be derived;

(2) Kits used, intended for use, or designed for use in manufacturing, compounding, converting, producing, processing, or preparing controlled substances;

(3) Isomerization devices used, intended for use, or designed for use in increasing the potency of any species of plant which is a controlled substance;

(4) Testing equipment used, intended for use, or designed for use in identifying or in analyzing the strength, effectiveness, or purity of controlled substances;

(5) Scales and balances used, intended for use, or designed for use in weighing or measuring controlled substances;

(6) Diluents and adulterants, such as quinine hydrochloride, mannitol, mannite, dextrose, and lactose, used, intended for use, or designed for use in cutting controlled substances;

(7) Separation gins and sifters used, intended for use, or designed for use in removing twigs and seeds from, or in otherwise cleaning or refining, marihuana;

(8) Blenders, bowls, containers, spoons, and mixing devices used, intended for use, or designed for use in compounding controlled substances;

(9) Capsules, balloons, envelopes, and other containers used, intended for use, or designed for use in packaging small quantities of controlled substances;

(10) Containers and other objects used, intended for use, or designed for use in storing or concealing controlled substances;

(11) Hypodermic syringes, needles, and other objects used, intended for use, or designed for use in parenterally injecting controlled substances into the human body;

(12) Objects used, intended for use, or designed for use in ingesting, inhaling, or otherwise introducing marihuana, cocaine, hashish, or hashish oil into the human body, such as:

(i) Metal, wooden, acrylic, glass, stone, plastic, or ceramic pipes with or without screens, permanent screens, hashish heads, or punctured metal bowls;
69.50.201 Title 69 RCW:  Food, Drugs, Cosmetics, and Poisons

(1) Enforcement of chapter—Authority to change schedules of controlled substances.  (a) The state board of pharmacy shall enforce this chapter and may add substances to or delete or reschedule substances listed in RCW 69.50.204, 69.50.206, 69.50.208, 69.50.210, or 69.50.212 pursuant to the procedures of chapter 34.05 RCW.

(b) After considering the factors enumerated in subsection (a) of this section, the board shall make findings with respect thereto and adopt and cause to be published a rule controlling the substance upon finding the substance has a potential for abuse.

(c) The board, without regard to the findings required by subsection (a) of this section or RCW 69.50.203, 69.50.205, 69.50.207, 69.50.209, and 69.50.211 or the procedures prescribed by subsections (a) and (b) of this section, may place an immediate precursor in the same schedule in which the controlled substance of which it is an immediate precursor is placed or in any other schedule. If the board designates a substance as an immediate precursor, substances that are precursors of the controlled precursor are not subject to control solely because they are precursors of the controlled precursor.

(d) If a substance is designated, rescheduled, or deleted as a controlled substance under federal law, the board shall similarly control the substance under this chapter after the expiration of thirty days from the date of publication in the federal register of a final order designating the substance as a controlled substance or rescheduling or deleting the substance or from the date of issuance of an order of temporary scheduling under Section 508 of the federal Dangerous Drug Diversion Control Act of 1984, 21 U.S.C. Sec. 811(h), unless within that thirty-day period, the board or an interested party objects to inclusion, rescheduling, or deletion. If no objection is made, the board shall adopt and cause to be published, without the necessity of making determinations or findings as required by subsection (a) of this section or RCW 69.50.203, 69.50.205, 69.50.207, 69.50.209, and 69.50.211, a final rule, for which notice of proposed rule making is omitted, designating, rescheduling, temporarily scheduling, or deleting the substance. If an objection is made, the board shall make a determination with respect to the designation, rescheduling, or deletion of the substance as provided by subsection (a) of this section. Upon receipt of an objection to inclusion, rescheduling, or deletion under this chapter by the board, the board shall publish notice of the
receipt of the objection, and control under this chapter is
stayed until the board adopts a rule as provided by subsection
(a) of this section.

(e) The board, by rule and without regard to the require-
ments of subsection (a) of this section, may schedule a sub-
stance in Schedule I regardless of whether the substance is
substantially similar to a controlled substance in Schedule I
or II if the board finds that scheduling of the substance on an
emergency basis is necessary to avoid an imminent hazard to
the public safety and the substance is not included in any
other schedule or no exemption or approval is in effect for the
substance under Section 505 of the federal Food, Drug, and
Cosmetic Act, 21 U.S.C. Sec. 355. Upon receipt of notice
under RCW 69.50.214, the board shall initiate scheduling of
the controlled substance analog on an emergency basis pursu-
ant to this subsection. The scheduling of a substance under
this subsection expires one year after the adoption of the
scheduling rule. With respect to the finding of an imminent
hazard to the public safety, the board shall consider whether
the substance has been scheduled on a temporary basis under
federal law or factors set forth in subsection (a)(1)(iv), (v),
and (vi) of this section, and may also consider clandestine
importation, manufacture, or distribution, and, if available,
information concerning the other factors set forth in subsec-
tion (a)(1) of this section. A rule may not be adopted under
this subsection until the board initiates a rule-making pro-
cedure under subsection (a) of this section with respect to
the substance. A rule adopted under this subsection must be
vacated upon the conclusion of the rule-making proceeding
initiated under subsection (a) of this section with respect to
the substance.

(g) [(f)] Authority to control under this section does not
extend to distilled spirits, wine, malt beverages, or tobacco as
those terms are defined or used in Titles 66 and 26 RCW.
[1998 c 245 § 108; 1993 c 187 § 2; 1989 1st ex.s. c 9 § 430;
1986 c 124 § 2; 1971 ex.s. c 308 § 69.50.201.]

Additional notes found at www.leg.wa.gov

69.50.202 Nomenclature. The controlled substances
listed or to be listed in the schedules in RCW 69.50.204,
69.50.206, 69.50.208, 69.50.210, and 69.50.212 are included
by whatever official, common, usual, chemical, or trade
name designated. [1971 ex.s. c 308 § 69.50.202.]

69.50.203 Schedule I tests. (a) The state board of phar-
macy shall place a substance in Schedule I upon finding that
the substance:

(1) has high potential for abuse;
(2) has no currently accepted medical use in treatment in the
United States; and
(3) lacks accepted safety for use in treatment under med-
cal supervision.

(b) The board may place a substance in Schedule I with-
out making the findings required by subsection (a) of this sec-
tion if the substance is controlled under Schedule I of the fed-
eral Controlled Substances Act by a federal agency as the
result of an international treaty, convention, or protocol.
[1993 c 187 § 3; 1971 ex.s. c 308 § 69.50.203.]

69.50.204 Schedule I. Unless specifically excepted by
state or federal law or regulation or more specifically
included in another schedule, the following controlled sub-
stances are listed in Schedule I:

(a) Any of the following opiates, including their isomers,
esters, ethers, salts, and salts of isomers, esters, and ethers
whenever the existence of these isomers, esters, ethers, and
salts is possible within the specific chemical designation:

(1) Acetyl-alpha-methylfentanyl (N-[1-(1-methyl-2-
phenethyl)-4-piperidinyl]-N-phenylacetamide);
(2) Acetylmethadol;
(3) Allylprodine;
(4) Alphacetylmethadol, except levo-alphacetylmethad-
adol, also known as levo-alpha-acetylmethadol, levometha-
dyl acetate, or LAAM;
(5) Alphameprodine;
(6) Alphamethadol;
(7) Alpha-methylfentanyl (N-[1-(alpha-methyl-beta-
ethyl) 4-piperidyl] propionanilide); (1-(1-methyl-2-
phenylethyl)-4-(N-propanilido) piperidine);
(8) Alpha-methylthiofentanyl (N-[1-methyl-2-(2-thie-
nyl)ethyl-4-piperidinyl]-N-phenylpropanamide);
(9) Benzethidine;
(10) Betacetylmethadol;
(11) Beta-hydroxyfentanyl (N-[1-(2-hydroxy-2-
phenethyl)-4-piperidinyl]-N-phenylpropanamide);
(12) Beta-hydroxy-3-methylfentanyl, some trade or
other names: N-[1-(2-hydroxy-2-phenethyl)-3-methyl-4-
piperidinyl]-N-phenylpropanamide;
(13) Betameprodine;
(14) Betamethadol;
(15) Betaprodine;
(16) Clonitazene;
(17) Dextromoramide;
(18) Diampromide;
(19) Diethylthiambutene;
(20) Difenoxin;
(21) Dimenoxadol;
(22) Dimepethanol;
(23) Dimethylthiambutene;
(24) Dioxaphetyl butyrate;
(25) Dipipanone;
(26) Ethylmethylthiambutene;
(27) Etonitazene;
(28) Etoperidine;
(29) Furethidine;
(30) Hydroxypethidine;
(31) Ketobemidone;
(32) Levomoramide;
(33) Levophenacylmorphan;
(34) 3-Methylfentanyl (N-[3-methyl-1-(2-phenylethyl)-
4-piperidinyl]-N-phenylpropionate);
(35) 3-Methylthiofentanyl (N-[3-methyl-1-(2-thi-
nyl)ethyl-4-piperidinyl]-N-phenylpropanamide);
(36) Morphetidine;
(37) MPPP (1-methyl-4-phenyl-4-propionoxy-piperi-
dine);
(38) Noracymethadol;
(39) Norlevorphanol;
(40) Normethadone;
(41) Norpipanone;
(42) Para-fluorofentanyl (N-(4-fluorophenyl)-N-[1-(2-
phenethyl)-4-piperidinyl] propanamide);
(43) PEPAP(1-(2-phenethyl)-4-phenyl-4-acetoxypiperidine);
(44) Phenadoxone;
(45) Phenampromide;
(46) Phenomorphan;
(47) Phenoperidine;
(48) Pir tramide;
(49) Proheptazine;
(50) Properidine;
(51) Propiram;
(52) Racemoramide;
(53) Thiofentanyl (N-phenyl-N-[1-(2-thienyl)ethyl]-4-piperidinyl)-propanamide);
(54) Tilidine;
(55) Trimeperidine.
(b) Opium derivatives. Unless specifically excepted or unless listed in another schedule, any of the following opium derivatives, including their salts, isomers, and salts of isomers whenever the existence of those salts, isomers, and salts of isomers is possible within the specific chemical designation:

1. Acetorphine;
2. Acetyldihydrocodeine;
3. Benzylmorphine;
4. Codeine methylbromide;
5. Codeine-N-Oxide;
6. Cyprenorphine;
7. Desomorphine;
8. Dihydromorphine;
9. Drotebanol;
10. Etorphine, except hydrochloride salt;
11. Heroin;
12. Hydromorphinol;
13. Methyldesorphine;
14. Methyldihydromorphine;
15. Morphine methylbromide;
16. Morphine methylsulfonate;
17. Morphine-N-Oxide;
18. Myrophine;
19. Nicocodeine;
20. Nicomorphine;
21. Normorphine;
22. Pholcodine;
23. Thebacon.
(c) Hallucinogenic substances. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following hallucinogenic substances, including their salts, isomers, and salts of isomers whenever the existence of those salts, isomers, and salts of isomers is possible within the specific chemical designation. For the purposes of this subsection only, the term "isomer" includes the optical, position, and geometric isomers:

1. Alpha-ethyltryptamine: Some trade or other names: Etryptamine; monase; a-ethyl-1H-indole-3-ethanamine; 3-(2-amino butyl) indole; a-ET; and AET;
2. 4-bromo-2,5-dimethoxyamphetamine: Some trade or other names: 4-bromo-2,5-dimethoxy-a-methylphenethylamine; 4-bromo-2,5-DMA;
3. 4-bromo-2,5-dimethoxyphenethylamine: Some trade or other names: 2-(4-bromo-2,5-dimethoxyphenyl)-1-aminoethane; alpha-desmethyl DOB; 2C-B; nexus;
4. 2,5-dimethoxyamphetamine: Some trade or other names: 2,5-dimethoxy-a-methylphenethylamine; 2,5-DMA;
5. 2,5-dimethoxy-4-ethylamphetamine (DOET);
6. 2,5-dimethoxy-4-(n)-propylthiophenethylamine:

Other names: 2C-T-7;
7. 4-methoxyamphetamine: Some trade or other names: 4-methoxy-a-methylphenethylamine; paramethoxyamphetamine, PMA;
8. 5-methoxy-3,4-methylenedioxyamphetamine;
9. 4-methyl-2,5-dimethoxyamphetamine: Some trade and other names: 4-methyl-2,5-dimethoxy-a-methylphenethylamine; "DOM"; and "STP";
10. 3,4-methylenedioxyamphetamine;
11. 3,4-methylenedioxymethamphetamine (MDMA);
12. 3,4-methylenedioxide-N-ethylamphetamine, also known as N-ethyl-alpha-methyl-3,4(methylenedioxy)phenethylamine, N-ethyl MDA, MDE, MDEA;
13. N-hydroxy-3,4-methylenedioxymethamphetamine also known as N-hydroxy-alpha-methyl-3,4(methylenedioxy)phenethylamine, N-hydroxy MDA;
14. 3,4,5-trimethoxyamphetamine;
15. Alpha-methyltryptamine: Other name: AMT;
16. Bufotenine: Some trade or other names: 3-(beta-Dimethylaminoethyl)-5-hydroxindole; 3-(2-dimethylaminoethyl)-5-indolol; N, N-dimethylserotonin; 5-hydroxy-N,N-dimethyltryptamine; mappine;
17. Diethyltryptamine: Some trade or other names: N,N-Diethyltryptamine; DET;
18. Dimethyltryptamine: Some trade or other names: DMT;
19. 5-methoxy-N,N-diisopropyltryptamine: Other name: 5-MeO-DIPT;
20. Ibogaine: Some trade or other names: 7-Ethyl-6,6 beta,7,8,9,10,12,13-octahydro-2-methoxy-6,9-methano-5H-pyrido(1'2',1,2) azezino (5,4-b) indole; Tabernanthe iboga;
21. Lysergic acid diethylamide;
22. Marihuana or marijuana;
23. Mescaline;
24. Parahexyl-7374: Some trade or other names: 3-Hexyl-1-hydroxy-7, 8, 9, 10-tetrahydro-6, 6, 9-trimethyl-6H-dibenzo[b,d]pyran; synhexyl;
25. Peyote, meaning all parts of the plant presently classified botanically as Lophophora Williamsii Lemaire, whether growing or not, the seeds thereof, any extract from any part of such plant, and every compound, manufacture, salts, derivative, mixture, or preparation of such plant, its seeds, or extracts; (interprets 21 U.S.C. Sec. 812 (c), Schedule I (c)(12));
26. N-ethyl-3-piperidyl benzilate; 
27. N-methyl-3-piperidyl benzilate; 
28. Psilocybin; 
29. Psilocyce; 
30. Tetrahydrocannabinols, meaning tetrahydrocannabinols naturally contained in a plant of the genus Cannabis (cannabis plant), as well as synthetic equivalents of the substances contained in the plant, or in the resinous extractives of Cannabis, species, and/or synthetic substances, derive-
tives, and their isomers with similar chemical structure and pharmacological activity such as the following:

(i) 1-cis- or trans tetrahydrocannabinol, and their optical isomers, excluding tetrahydrocannabinol in sesame oil and encapsulated in a soft gelatin capsule in a drug product approved by the United States Food and Drug Administration;

(ii) 6-cis- or trans tetrahydrocannabinol, and their optical isomers;

(iii) 3,4-cis- or trans tetrahydrocannabinol, and its optical isomers;

(Since nomenclature of these substances is not internationally standardized, compounds of these structures, regardless of numerical designation of atomic positions covered.)

(31) Ethylamine analog of phencyclidine: Some trade or other names: N-ethyl-1-phenylethylamine, (1-phenethyl)cyclohexyl ethylamine; N-(1-phenylethyl)cyclohexyl ethylamine; cyclohexylamine; PCE;

(32) Pyrrolidine analog of phencyclidine: Some trade or other names: 1-(1-phenylcyclohexyl)piperidine; PCPy; PHP;

(33) Thiophene analog of phencyclidine: Some trade or other names: 1-(1-[2-thienyl]-cyclohexyl)-pipendine; 2-thienyl analog of phencyclidine; TPCP; TCP;

(34) 1-[1-(2-thienyl)cyclohexyl]piperidine: A trade or other name is TCPy.

(d) Depressants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances having a depressant effect on the central nervous system, including its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation.

(1) Gamma-hydroxybutyric acid: Some other names include GHB; gamma-hydroxybutyrate; 4-hydroxybutyrate; 4-hydroxybutanoic acid; sodium oxybate; sodium oxybutyraye;

(2) Mecloqualone;

(3) Methaqualone.

(e) Stimulants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances having a stimulant effect on the central nervous system, including its salts, isomers, and salts of isomers.

(1) Aminorex: Some other names: aminoxaphen; 2-amino-5-phenyl-2-oxazoline; or 4, 5-dihydro-5-phenyl-2-oxazolamine;

(2) N-Benzylpiperazin: Some other names: BZP,1-benzylpiperazin;

(3) Cathinone, also known as 2-amino-1-phenyl-1-propa
none, alpha-aminopropiophenone, 2-aminopropiophenone and norephedrone;

(4) Fenethylline;

(5) Methanthionine: Some other names: 2-(methylamino)-propiophenone; alpha-(methylamino)propiophenone; 2-(methylamino)-1-phenylpropan-1-one; alpha-N-methylaminopropiophenone; monomethylpropion; ephedrine; N-methylcathinone; methylcathinone; AL-464; AL-422; AL-463 and UR1432, its salts, optical isomers, and salts of optical isomers;

(6) (+)-cis-4-methylaminorex ((+)-cis-4,5-dihydro-4-methyl-5-phenyl-2-oxazololine);
(xviii) Thebaine.

(2) Any salt, compound, isomer, derivative, or preparation thereof that is chemically equivalent or identical with any of the substances referred to in subsection (b)(1) of this section, but not including the isoquinoline alkaloids of opium.

(3) Opium poppy and poppy straw.

(4) Coca leaves and any salt, compound, derivative, or preparation of coca leaves including cocaine and ecorpine, and their salts, isomers, derivatives, and salts of isomers and derivatives, and any salt, compound, derivative, or preparation thereof which is chemically equivalent or identical with any of these substances, except that the substances shall not include decocainized coca leaves or extractions of coca leaves which do not contain cocaine or ecorpine.

(5) Concentrate of poppy straw (The crude extract of poppy straw in either liquid, solid, or powder form which contains the phenanthrene alkaloids of the opium poppy.)

(c) Opiates. Unless specifically excepted or unless in another schedule, any of the following synthetic opiates, including its isomers, esters, ethers, salts, and salts of isomers, esters, and ethers, whenever the existence of such isomers, esters, ethers, and salts is possible within the specific chemical designation, dextrophan and levopropoxyphene excepted:

(1) Alfentanil;
(2) Alphaprodine;
(3) Anileridine;
(4) Bezitramide;
(5) Bulk dextropropoxyphene (nondosage forms);
(6) Carfentanil;
(7) Dihydrocodeine;
(8) Diphenoxylate;
(9) Fentanyl;
(10) Isomethadone;
(11) Levo-alpha-acetyl-methadol, also known as levomethadyl acetate, or LAAM;
(12) Levo-methadone;
(13) Levorphanol;
(14) Metazocine;
(15) Methadone;
(16) Methadone—Intermediate, 4-cyano-2-dimethylamino-4, 4-diphenyl butanol;
(17) Moramide—Intermediate, 2-methyl-3-morpholinol1, 1-diphenylpropane-carboxylic acid;
(18) Pethidine (meperidine);
(19) Pethidine—Intermediate-A, 4-cyano-1-methyl-4-phenylpipеридине;
(20) Pethidine—Intermediate-B, ethyl-4-phenylpipеридине-4-carboxylate;
(21) Pethidine—Intermediate-C, 1-methyl-4-phenylpipеридине-4-carboxylic acid;
(22) Phenazocine;
(23) Pimino-dine;
(24) Racemethorphan;
(25) Racemorphan;
(26) Remifentanil;
(27) Sufentanil;
(28) Tapentadol.

(d) Stimulants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances having a stimulant effect on the central nervous system:

(1) Amphetamine, its salts, optical isomers, and salts of its optical isomers;
(2) Methamphetamine, its salts, isomers, and salts of its isomers;
(3) Phenmetrazine and its salts;
(4) Methylphenidate;
(5) Lisdexamfetamine, its salts, isomers, and salts of its isomers.

(e) Depressants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances having a depressant effect on the central nervous system, including its salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation:

(1) Amobarbital;
(2) Glutethimide;
(3) Pentobarbital;
(4) Phencyclidine;
(5) Secobarbital.

(f) Hallucinogenic substances.

Nabilone: Some trade or other names are (±)-trans-3-(1,1-dimethylheptyl)-6,6a,7,8,10a-hexahydro-1-hydroxy-6,6-dimethyl-9H-dibenzo[b,d]pyran-9-one.

(g) Immediate precursors. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances:

(1) Immediate precursor to amphetamine and methamphetamine:

(i) Phenylacetone: Some trade or other names phenyl-2-propanone, P2P, benzyl methyl ketone, methyl benzyl ketone.

(2) Immediate precursors to phencyclidine (PCP):

(i) 1-phenylcyclohexylamine;
(ii) 1-piperidinocyclohexancarbonitrile (PCC).

The controlled substances in this section may be added, rescheduled, or deleted as provided for in RCW 69.50.201. [2010 c 177 § 3; 1993 c 187 § 6; 1986 c 124 § 4; 1980 c 138 § 2; 1971 ex.s. c 308 § 69.50.206.]

State board of pharmacy may change schedules of controlled substances: RCW 69.50.201.

69.50.207 Schedule III tests. (a) The state board of pharmacy shall place a substance in Schedule III upon finding that:

(1) the substance has a potential for abuse less than the substances included in Schedules I and II;

(2) the substance has currently accepted medical use in treatment in the United States; and

(3) abuse of the substance may lead to moderate or low physical dependence or high psychological dependence.

(b) The state board of pharmacy may place a substance in Schedule III without making the findings required by subsection (a) of this section if the substance is controlled under Schedule III of the federal Controlled Substances Act by a federal agency as the result of an international treaty, conven-
69.50.208 Schedule III. Unless specifically excepted by state or federal law or regulation or more specifically included in another schedule, the following controlled substances are listed in Schedule III:

(a) Stimulants. Any material, compound, mixture, or preparation containing any quantity of the following substances having a stimulant effect on the central nervous system, including their salts, isomers, whether optical, position, or geometric, and salts of isomers whenever the existence of those salts, isomers, and salts of isomers is possible within the specific chemical designation:

(1) Any compound, mixture, or preparation in dosage unit form containing any stimulant substance included in Schedule II and which was listed as an excepted compound on August 25, 1971, pursuant to the federal Controlled Substances Act, and any other drug of the quantitative composition shown in that list for those drugs or which is the same except for containing a lesser quantity of controlled substances:

(2) Benzphetamine;
(3) Chlorphentermine;
(4) Clortermine;
(5) Methamphetamine;

(b) Depressants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances having a depressant effect on the central nervous system:

(1) Any compound, mixture, or preparation containing:

(i) Amobarbital;
(ii) Secobarbital;
(iii) Pentobarbital;

or any salt thereof and one or more other active medicinal ingredients which are not listed in any schedule;

(2) Any suppository dosage form containing:

(i) Amobarbital;
(ii) Secobarbital;
(iii) Pentobarbital;

or any salt of any of these drugs and approved by the Food and Drug Administration for marketing only as a suppository;

(3) Any substance which contains any quantity of a derivative of barbituric acid, or any salt of a derivative of barbituric acid;

(4) Chlorhexadol;
(5) Embutramide;

(6) Any drug product containing gamma hydroxybutyric acid, including its salts, isomers, and salts of isomers, for which an application is approved under section 505 of the federal food, drug, and cosmetic act;

(7) Ketamine, its salts, isomers, and salts of isomers, some other names for ketamine: (<plus-minus>)-2-(2-chlorophenyl)-2-(methylamino)-cyclohexanone;

(8) Lysergic acid;
(9) Lysergic acid amide;
(10) Methyprylon;
(11) Sulfondiethylmethylene;
(12) Sulfonethylmethylene;

(13) Sulfonmethane;
(14) Tiletamine and zolazepam or any of their salts—some trade or other names for a tiletamine-zolazepam combination product: Telazol, some trade or other names for tiletamine: 2-(ethylamino)-2-(2-thienyl) cyclohexanone, some trade or other names for zolazepam: 4-(2-fluorophenyl)-6,8-dihydro-1,3,8-trimethylpyrazolo-[3,4-e][1,4]-diazepin-7(1H)-one flupryrazapam.

(c) Nalorphine.

(d) Narcotic drugs. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation containing limited quantities of any of the following narcotic drugs, or any salts thereof calculated as the free anhydrous base or alkaloid, in limited quantities as set forth in this subsection:

(1) Not more than 1.8 grams of codeine per 100 milliliters or not more than 90 milligrams per dosage unit, with an equal or greater quantity of an isoquinoline alkaloid of opium;

(2) Not more than 1.8 grams of codeine per 100 milliliters or not more than 90 milligrams per dosage unit, with one or more active, nonnarcotic ingredients in recognized therapeutic amounts;

(3) Not more than 300 milligrams of dihydrocodeine (hydrocodone) per 100 milliliters or not more than 15 milligrams per dosage unit, with a fourfold or greater quantity of an isoquinoline alkaloid of opium;

(4) Not more than 300 milligrams of dihydrocodeine (hydrocodone) per 100 milliliters or not more than 15 milligrams per dosage unit, with one or more active, nonnarcotic ingredients in recognized therapeutic amounts;

(5) Not more than 1.8 grams of dihydrocodeine per 100 milliliters or not more than 90 milligrams per dosage unit, with one or more active, nonnarcotic ingredients in recognized therapeutic amounts;

(6) Not more than 300 milligrams of ethylmorphine per 100 milliliters or not more than 15 milligrams per dosage unit, with one or more active, nonnarcotic ingredients in recognized therapeutic amounts;

(7) Not more than 500 milligrams of opium per 100 milliliters or per 100 grams, or not more than 25 milligrams per dosage unit, with one or more active, nonnarcotic ingredients in recognized therapeutic amounts; and

(8) Not more than 50 milligrams of morphine per 100 milliliters or per 100 grams with one or more active, nonnarcotic ingredients in recognized therapeutic amounts.

(e) Any material, compound, mixture, or preparation containing any of the following narcotic drugs or their salts: Buprenorphine.

(f) Hallucinogenic substances. Dronabinol (synthetic) in sesame oil and encapsulated in a soft gelatin capsule in a United States food and drug administration approved product. Some other names for dronabinol: [6a-R-trans]-6a,7,8,10a-tetrahydro-6,6,9-trimethyl-3-pentyl-6H-dibenzo[b,d]pyran-1-ol, or (-)-delta-9-(trans)-tetrahydrocannabinol.

(g) Anabolic steroids. The term "anabolic steroids" means any drug or hormonal substance, chemically and pharmacologically related to testosterone, other than estrogens, progestins, corticosteroids, and dehydroepiandrosterone, that promotes muscle growth and includes:

1. 3β,17-dihydroxy-5α-androstane;
(2) 3α,17β-dihydroxy-5α-androstan-3-one;
(3) 5α-androstan-3,17-dione;
(4) 1-androstenediol (3β,17β-dihydroxy-5α-androst-1-en-3-one);
(5) 1-androstenediol (3α,17β-dihydroxy-5α-androst-1-en-3-one);
(6) 4-androstenediol (3β,17β-dihydroxy-androst-4-en-3-one);
(7) 5-androstenediol (3β,17β-dihydroxy-androst-5-en-3-one);
(8) 1-androstenedione (5α-androst-1-en-3,17-dione);
(9) 4-androstenedione (androst-4-en-3,17-dione);
(10) 5-androstenedione (androst-5-en-3,17-dione);
(11) Bolasterone (7α,17α-dimethyl-17β-hydroxyandrost-4-en-3-one);
(12) Boldenone (17β-hydroxyandrost-1,4-diene-3-one);
(13) Calusterone (7β,17α-dimethyl-17β-hydroxyandrost-4-en-3-one);
(14) Clostebol (4-chloro-17β-hydroxyandrost-4-en-3-one);
(15) Dehydrochloromethyltestosterone (4-chloro-17β-hydroxy-17α-methyl-androst-1,4-dien-3-one);
(16) Δ1-dihydrotestosterone (a.k.a. '1-testosterone')
(17) 4-dihydrotestosterone (17β-hydroxy-androst-3-en-3-one);
(18) Drostanolone (17β-hydroxy-2α-methyl-5α-androst-3-en-3-one);
(19) Ethylestrenol (17α-ethyl-17β-hydroxyestr-4-en-3-one);
(20) Fluoxymesterone (9-fluoro-17α-methyl-11β,17γ-dihydroxyandrost-4-en-3-one);
(21) Formebolone (2-formyl-17α-methyl-11β,17δ-dihydroxyandrost-1,4-dien-3-one);
(22) Furazabol (17α-methyl-17β-hydroxyandrostan[2,3-c]-furazan);
(23) 13β-ethyl-17β-hydroxyestra-4,9,11-trien-3-one;
(24) 4-hydroxytestosterone (4,17β-dihydroxy-androst-4-en-3-one);
(25) 4-hydroxy-19-nortestosterone (4,17β-dihydroxyestr-4-en-3-one);
(26) Mesterolone (17α-methyl-17β-hydroxy-5-androst-3-en-3-one);
(27) Mesterolone (1α-methyl-17β-hydroxy-[5α]-androstan-3-one);
(28) Methandienone (17α-methyl-17β-hydroxyandrost-1,4-dien-3-one);
(29) Methandriol (17α-methyl-3β,17β-dihydroxyandrost-5-en-3-one);
(30) Methenolone (1-methyl-17β-hydroxy-5α-androst-1-en-3-one);
(31) 17α-methyl-3β,17β-dihydroxy-5α-androstane;
(32) 17α-methyl-5α,17β-dihydroxy-5α-androstane;
(33) 17α-methyl-3β,17β-dihydroxyandrost-4-en-3-one;
(34) 17α-methyl-4-hydroxyandrolone (17α-methyl-4-hydroxy-17β-hydroxyestr-4-en-3-one);
(35) Methyldienolone (17α-methyl-17β-hydroxyestr-4,9(10)-dien-3-one);
(36) Methyltrienolone (17α-methyl-17β-hydroxyestr-4,9,11-trien-3-one);
(37) Methyltestosterone (17α-methyl-17β-hydroxyandrost-4-en-3-one);
(38) Mibolerone (7α,17α-dimethyl-17β-hydroxyestr-4-en-3-one);
(39) 17α-methyl-17β-hydroxy-17α-methyl-5α-androst-1-en-3-one (also known as '17α-methyl-1-testosterone');
(40) Nandrolone (17β-hydroxyestr-4-en-3-one);
(41) 19-nor-4-androstenediol (3β,17β-dihydroxyestr-4-en-3-one);
(42) 19-nor-4-androstenediol (3α,17β-dihydroxyestr-4-en-3-one);
(43) 19-nor-5-androstenediol (3β,17β-dihydroxyestr-5-en-3-one);
(44) 19-nor-5-androstenediol (3α,17β-dihydroxyestr-5-en-3-one);
(45) 19-nor-4-androstenedione (estr-4-en-3,17-dione);
(46) 19-nor-5-androstenedione (estr-5-en-3,17-dione);
(47) Norbolethone (13β, 17α-diethyl-17β-hydroxygon-4-en-3-one);
(48) Norclostebol (4-chloro-17β-hydroxyestr-4-en-3-one);
(49) Norethandrolone (17α-ethyl-17β-hydroxyestr-4-en-3-one);
(50) Normethandrolone (17α-methyl-17β-hydroxyestr-4-en-3-one);
(51) Oxandrolone (17α-methyl-17β-hydroxy-2-oxa-[5α]-androstan-3-one);
(52) Oxymesterone (17α-methyl-4,17β-dihydroxyandrost-4-en-3-one);
(53) Oxymetholone (17α-methyl-2-hydroxymethylene-17β-hydroxy-[5α]-androstan-3-one);
(54) Stanozolol (17α-methyl-17β-hydroxy-[5α]-androstan-2-en[3,2-c]-pyrazole);
(55) Stenbolone (17β-hydroxy-2-methyl-[5α]-androstan-1-en-3-one);
(56) Testolactone (13-hydroxy-3-oxo-13,17-secoandrosta-1,4-dien-17-ic acid lactone);
(57) Testosterone (17β-hydroxyandrost-4-en-3-one);
(58) Tetrahydrogestrinone (13β, 17α-diethyl-17β-hydroxygon-4,9,11-trien-3-one);
(59) Trenbolone (17β-hydroxyestr-4,9,11-trien-3-one); and
(60) Any salt, ester, or ether of a drug or substance described in this section. Such term does not include an anabolic steroid that is expressly intended for administration through implants to cattle or other nonhuman species and that has been approved by the secretary of the department of health and human services for such administration. If any person prescribes, dispenses, or distributes such steroid for human use, the person shall be considered to have prescribed, dispensed, or distributed an anabolic steroid within the meaning of this section.

The state board of pharmacy may except by rule any compound, mixture, or preparation containing any stimulant or depressant substance listed in subsection (a)(1) and (2) of this section from the application of all or any part of this chapter if the compound, mixture, or preparation contains

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69.50.209 Schedule IV tests. (a) The state board of pharmacy shall place a substance in Schedule IV upon finding that:

(1) the substance has a low potential for abuse relative to substances in Schedule III;
(2) the substance has currently accepted medical use in treatment in the United States; and
(3) abuse of the substance may lead to limited physical dependence or psychological dependence relative to the substances included in Schedule III.

(b) The state board of pharmacy may place a substance in Schedule IV without making the findings required by subsection (a) of this section if the substance is controlled under Schedule IV of the federal Controlled Substances Act by a federal agency as the result of an international treaty, convention, or protocol. [1993 c 187 § 9; 1971 ex.s. c 308 § 69.50.209.]

69.50.210 Schedule IV. Unless specifically excepted by state or federal law or regulation or more specifically included in another schedule, the following controlled substances are listed in Schedule IV:

(a) Any material, compound, mixture, or preparation containing any of the following narcotic drugs, or their salts calculated as the free anhydrous base or alkaloid, in limited quantities as set forth below:

(1) Not more than 1 milligram of difenoxin and not less than 25 micrograms of atropine sulfate per dosage unit.
(2) Dextropropoxyphene (alpha-(+)-4-dimethylamino-1,2-diphenyl-3-methyl-2-propionoxybutane).

(b) Depressants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation containing any quantity of the following substances having a depressant effect on the central nervous system, including their salts, isomers, and salts of isomers whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation:

1. Alprazolam;
2. Barbital;
3. Bromazepam;
4. Camazepam;
5. Carisoprodol;
6. Chloral betaine;
7. Chloral hydrate;
8. Chlordiazepoxide;
9. Clobazam;
10. Clonazepam;
11. Clorazepate;
12. Clotiazepam;
13. Cloxazolam;
14. Delorazepam;
15. Diazepam;
16. Dichloralphenazone;
17. Estazolam;
18. Ethchlorvynol;
19. Ethinamate;
20. Ethyl loflazepate;
21. Fludiazepam;
22. Flunitrazepam;
23. Flurazepam;
24. Halazepam;
25. Haloxazolam;
26. Ketazolam;
27. Loprazolam;
28. Lorazepam;
29. Lormetazepam;
30. Mebutamate;
31. Medazepam;
32. Meprobamate;
33. Methohexital;
34. Methylphenobarbital (mephobarbital);
35. Midazolam;
36. Nimetazepam;
37. Nitrazepam;
38. Nordiazepam;
39. Oxazepam;
40. Oxazolam;
41. Paraldehyde;
42. Petrichloral;
43. Phenobarbital;
44. Pinazepam;
45. Prazepam;
46. Quazepam;
47. Temazepam;
48. Tetrazepam;
49. Triazolam;
50. Zaleplon;
51. Zolpidem; and
52. Zopiclone.

(c) Fenfluramine. Any material, compound, mixture, or preparation containing any quantity of the following substance, including its salts, isomers, and salts of such isomers, whenever the existence of such salts, isomers, and salts of isomers is possible: Fenfluramine.

(d) Stimulants. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation containing any quantity of the following substances having a stimulant effect on the central nervous system, including their salts, isomers, and salts of isomers:

1. Cathine((+)norpseudoephedrine);
2. Diethylpropion;
3. Fenfluramine;
4. Fenproporex;
5. Mazindol;
6. Mefenorex;
7. Modafinil;
8. Pemoline (including organometallic complexes and chelates thereof);
9. Phentermine;
(10) Pipradrol;
(11) Sibutramine;
(12) SPA ((-)-1-dimethylamino-1, 2-dephenylethane).

(e) Other substances. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation containing any quantity of the following substance, including its salts:

(1) Pentazocine;
(2) Butorphanol, including its optical isomers.

The state board of pharmacy may except by rule any compound, mixture, or preparation containing any depressant substance listed in subsection (b) of this section from the application of all or any part of this chapter if the compound, mixture, or preparation contains one or more active medicinal ingredients not having a depressant effect on the central nervous system, and if the admixtures are in combinations, quantity, proportion, or concentration that vitiate the potential for abuse of the substances having a depressant effect on the central nervous system.

The controlled substances listed in this section may be added, rescinded, or deleted as provided for in RCW 69.50.201. [2010 c 177 § 5; 1993 c 187 § 10; 1986 c 124 § 6; 1981 c 147 § 2; 1980 c 138 § 4; 1971 ex.s. c 308 § 69.50.210.]

State board of pharmacy may change schedules of controlled substances: RCW 69.50.201.

69.50.211 Schedule V tests. (a) The state board of pharmacy shall place a substance in Schedule V upon finding that:

(1) the substance has low potential for abuse relative to the controlled substances included in Schedule IV;
(2) the substance has currently accepted medical use in treatment in the United States; and
(3) abuse of the substance may lead to limited physical dependence or psychological dependence relative to the substances included in Schedule IV.

(b) The state board of pharmacy may place a substance in Schedule V without being required to make the findings required by subsection (a) of this section if the substance is controlled under Schedule V of the federal Controlled Substances Act by a federal agency as the result of an international treaty, convention, or protocol. [1993 c 187 § 11; 1971 ex.s. c 308 § 69.50.211.]

69.50.212 Schedule V. Unless specifically excepted by state or federal law or regulation or more specifically included in another schedule, the following controlled substances are listed in Schedule V:

(a) Any compound, mixture, or preparation containing any of the following narcotic drugs, or their salts calculated as the free anhydrous base or alkaloid, in limited quantities as set forth in this subsection, which also contains one or more nonnarcotic active medicinal ingredients in sufficient proportion to confer upon the compound, mixture, or preparation, valuable medicinal qualities other than those possessed by the narcotic drug alone:

(1) Not more than 200 milligrams of codeine per 100 milliliters or per 100 grams;
(2) Not more than 100 milligrams of dihydrocodeine per 100 milliliters or per 100 grams;
(3) Not more than 100 milligrams of ethylmorphine per 100 milliliters or per 100 grams;
(4) Not more than 2.5 milligrams of diphenoxylate and not less than 25 micrograms of atropine sulfate per dosage unit;
(5) Not more than 100 milligrams of opium per 100 milliliters or per 100 grams;
(6) Not more than 0.5 milligrams of difenoxin and not less than 25 micrograms of atropine sulfate per dosage unit.

(b) Stimulants. Unless specifically exempted or excluded or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances having a stimulant effect on the central nervous system, including its salts, isomers, and salts of isomers: Pyrovalerone.

(c) Depressants. Unless specifically exempted or excluded or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substances having a depressant effect on the central nervous system, including its salts:

[Racemic][(-)-2-acetoamido-N-benzyl-3-methoxy-propionamide];
(2) Pregabalin[(S)-3-(aminomethyl)-5-methylhexanoic acid].

The state board of pharmacy shall place a substance in Schedule V upon finding that:

(1) the substance is included in another schedule, the following controlled substances or more specifically:
(2) the substance has currently accepted medical use in treatment in the United States; and
(3) abuse of the substance may lead to limited physical dependence or psychological dependence relative to the substances included in Schedule IV.

The controlled substances listed in this section may be added, rescinded, or deleted as provided for in RCW 69.50.201. [2010 c 177 § 6; 1993 c 187 § 12; 1986 c 124 § 7; 1980 c 138 § 5; 1971 ex.s. c 308 § 69.50.212.]

State board of pharmacy may change schedules of controlled substances: RCW 69.50.201.

69.50.213 Republishing of schedules. The state board of pharmacy shall publish updated schedules annually. Failure to publish updated schedules is not a defense in any administrative or judicial proceeding under this chapter. [1993 c 187 § 13; 1971 ex.s. c 308 § 69.50.213.]

69.50.214 Controlled substance analog. A controlled substance analog, to the extent intended for human consumption, shall be treated, for the purposes of this chapter, as a substance included in Schedule I. Within thirty days after the initiation of prosecution with respect to a controlled substance analog by indictment or information, the prosecuting attorney shall notify the state board of pharmacy of information relevant to emergency scheduling as provided for in *RCW 69.50.201(f). After final determination that the controlled substance analog should not be scheduled, no prosecution relating to that substance as a controlled substance analog may continue or take place. [1993 c 187 § 14.]

*Reviser’s note: RCW 69.50.201 was amended by 1998 c 245 § 108, changing subsection (f) to subsection (e).
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69.50.302 Registration requirements. (a) Every person who manufactures, distributes, or dispenses any controlled substance within this state or who proposes to engage in the manufacture, distribution, or dispensing of any controlled substance within this state, shall obtain annually a registration issued by the department in accordance with the board’s rules.

(b) A person registered by the department under this chapter to manufacture, distribute, dispense, or conduct research with controlled substances may possess, manufacture, distribute, dispense, or conduct research with those substances to the extent authorized by the registration and in conformity with this Article.

(c) The following persons need not register and may lawfully possess controlled substances under this chapter:

(1) An agent or employee of any registered manufacturer, distributor, or dispenser of any controlled substance if the agent or employee is acting in the usual course of business or employment. This exemption shall not include any agent or employee distributing sample controlled substances to practitioners without an order;

(2) A common or contract carrier or warehouse operator, or an employee thereof, whose possession of any controlled substance is in the usual course of business or employment;

(3) An ultimate user or a person in possession of any controlled substance pursuant to a lawful order of a practitioner or in lawful possession of a substance included in Schedule V.

(d) The board may waive by rule the requirement for registration of certain manufacturers, distributors, or dispensers upon finding it consistent with the public health and safety. Personal practitioners licensed or registered in the state of Washington under the respective professional licensing acts shall not be required to be registered under this chapter unless the specific exemption is denied pursuant to RCW 69.50.305 for violation of any provisions of this chapter.

(e) A separate registration is required at each principal place of business or professional practice where the applicant manufactures, distributes, or dispenses controlled substances.

(f) The department may inspect the establishment of a registrant or applicant for registration in accordance with rules adopted by the board. [2011 c 336 § 839; 1993 c 187 § 16; 1989 1st ex.s. c 9 § 432; 1971 ex.s. c 308 § 69.50.302.]

Additional notes found at www.leg.wa.gov

69.50.303 Registration. (a) The department shall register an applicant to manufacture or distribute controlled substances included in RCW 69.50.204, 69.50.206, 69.50.208, 69.50.210, and 69.50.212 unless the board determines that the issuance of that registration would be inconsistent with the public interest. In determining the public interest, the board shall consider the following factors:

(1) maintenance of effective controls against diversion of controlled substances into other than legitimate medical, scientific, research, or industrial channels;

(2) compliance with applicable state and local law;

(3) promotion of technical advances in the art of manufacturing controlled substances and the development of new substances;

(4) any convictions of the applicant under any laws of another country or federal or state laws relating to any controlled substance;

(5) past experience in the manufacture or distribution of controlled substances, and the existence in the applicant’s establishment of effective controls against diversion of controlled substances into other than legitimate medical, scientific, research, or industrial channels;

(6) furnishing by the applicant of false or fraudulent material in any application filed under this chapter;

(7) suspension or revocation of the applicant’s federal registration to manufacture, distribute, or dispense controlled substances as authorized by federal law; and

(8) any other factors relevant to and consistent with the public health and safety.

(b) Registration under subsection (a) of this section does not entitle a registrant to manufacture or distribute controlled substances included in Schedule I or II other than those specified in the registration.

(c) Practitioners must be registered, or exempted under RCW 69.50.302(d), to dispense any controlled substances or to conduct research with controlled substances included in Schedules II through V if they are authorized to dispense or conduct research under the law of this state. The board need not require separate registration under this Article for practitioners engaging in research with nonnarcotic substances included in Schedules II through V where the registrant is already registered under this Article in another capacity. Practitioners registered under federal law to conduct research with substances included in Schedule I may conduct research with substances included in Schedule I within this state upon furnishing the board evidence of that federal registration.

(d) A manufacturer or distributor registered under the federal Controlled Substances Act, 21 U.S.C. Sec. 801 et seq., may submit a copy of the federal application as an application for registration as a manufacturer or distributor under this section. The board may require a manufacturer or distributor to submit information in addition to the application for registration under the federal act. [1993 c 187 § 17; 1989 1st ex.s. c 9 § 432; 1971 ex.s. c 308 § 69.50.303.]

Additional notes found at www.leg.wa.gov

69.50.304 Revocation and suspension of registration—Seizure or placement under seal of controlled substances. (a) A registration, or exemption from registration, under RCW 69.50.303 to manufacture, distribute, or dispense a controlled substance may be suspended or revoked by the state board of pharmacy upon finding that the registrant has:

(1) furnished false or fraudulent material information in any application filed under this chapter;

(2) been convicted of a felony under any state or federal law relating to any controlled substance;

(3) had the registrant’s federal registration suspended or revoked and is no longer authorized by federal law to manufacture, distribute, or dispense controlled substances; or

(4) committed acts that would render registration under RCW 69.50.303 inconsistent with the public interest as determined under that section.

Additional notes found at www.leg.wa.gov
(b) The board may limit revocation or suspension of a registration to the particular controlled substance or schedule of controlled substances, with respect to which grounds for revocation or suspension exist.

(c) If the board suspends or revokes a registration, all controlled substances owned or possessed by the registrant at the time of suspension or the effective date of the revocation order may be placed under seal. No disposition may be made of substances under seal until the time for taking an appeal has elapsed or until all appeals have been concluded unless a court, upon application, orders the sale of perishable substances and the deposit of the proceeds of the sale with the court. Upon a revocation order becoming final, all controlled substances may be forfeited to the state.

(d) The department may seize or place under seal any controlled substance owned or possessed by a registrant whose registration has expired or who has ceased to practice or do business in the manner contemplated by the registration. The controlled substance must be held for the benefit of the registrant or the registrant’s successor in interest. The department shall notify a registrant, or the registrant’s successor in interest, who has any controlled substance seized or placed under seal, of the procedures to be followed to secure the return of the controlled substance and the conditions under which it will be returned. The department may not dispose of any controlled substance seized or placed under seal under this subsection until the expiration of one hundred eighty days after the controlled substance was seized or placed under seal. The costs incurred by the department in seizing, placing under seal, maintaining custody, and disposing of any controlled substance under this subsection may be recovered from the registrant, any proceeds obtained from the disposition of the controlled substance, or from both. Any balance remaining after the costs have been recovered from the proceeds of any disposition must be delivered to the registrant or the registrant’s successor in interest.

(e) The department shall promptly notify the drug enforcement administration of all orders restricting, suspending, or revoking registration and all forfeitures of controlled substances. [1993 c 187 § 18; 1989 1st ex.s. c 9 § 434; 1986 c 124 § 8; 1971 ex.s. c 308 § 69.50.304.]

Additional notes found at www.leg.wa.gov

69.50.305 Procedure for denial, suspension, or revocation of registration. (a) Any registration, or exemption from registration, issued pursuant to the provisions of this chapter shall not be denied, suspended, or revoked unless the board denies, suspends, or revokes such registration, or exemption from registration, by proceedings consistent with the administrative procedure act, chapter 34.05 RCW.

(b) The board may suspend any registration simultaneously with the institution of proceedings under RCW 69.50.304, or where renewal of registration is refused, if it finds that there is an imminent danger to the public health or safety which warrants this action. The suspension shall continue in effect until the conclusion of the proceedings, including judicial review thereof, unless sooner withdrawn by the board or dissolving by a court of competent jurisdiction. [1971 ex.s. c 308 § 69.50.305.]

69.50.306 Records of registrants. Persons registered, or exempted from registration under RCW 69.50.302(d), to manufacture, distribute, dispense, or administer controlled substances under this chapter shall keep records and maintain inventories in conformance with the recordkeeping and inventory requirements of federal law and with any additional rules the state board of pharmacy issues. [1971 ex.s. c 308 § 69.50.306.]

69.50.308 Prescriptions. (a) A controlled substance may be dispensed only as provided in this section.

(b) Except when dispensed directly by a practitioner authorized to prescribe or administer a controlled substance, other than a pharmacy, to an ultimate user, a substance included in Schedule II may not be dispensed without the written prescription of a practitioner.

(1) Schedule II narcotic substances may be dispensed by a pharmacy pursuant to a facsimile prescription under the following circumstances:

(i) The facsimile prescription is transmitted by a practitioner to the pharmacy; and

(ii) The facsimile prescription is for a patient in a long-term care facility. "Long-term care facility" means nursing homes licensed under chapter 18.51 RCW, assisted living facilities licensed under chapter 18.20 RCW, and adult family homes licensed under chapter 70.128 RCW; or

(iii) The facsimile prescription is for a patient of a hospice program certified or paid for by medicare under Title XVIII; or

(iv) The facsimile prescription is for a patient of a hospice program licensed by the state; and

(v) The practitioner or the practitioner’s agent notes on the facsimile prescription that the patient is a long-term care or hospice patient.

(2) Injectable Schedule II narcotic substances that are to be compounded for patient use may be dispensed by a pharmacy pursuant to a facsimile prescription if the facsimile prescription is transmitted by a practitioner to the pharmacy.

(3) Under (1) and (2) of this subsection the facsimile prescription shall serve as the original prescription and shall be maintained as other Schedule II narcotic substances prescriptions.

(c) In emergency situations, as defined by rule of the state board of pharmacy, a substance included in Schedule II may be dispensed upon oral prescription of a practitioner, reduced promptly to writing and filed by the pharmacy. Prescriptions shall be retained in conformance with the requirements of RCW 69.50.306. A prescription for a substance included in Schedule II may not be refilled.

(d) Except when dispensed directly by a practitioner authorized to prescribe or administer a controlled substance, other than a pharmacy, to an ultimate user, a substance included in Schedule III or IV, which is a prescription drug as determined under RCW 69.04.560, may not be dispensed without a written or oral prescription of a practitioner. Any oral prescription must be promptly reduced to writing. The prescription shall not be filled or refilled more than six months after the date thereof or be refilled more than five times, unless renewed by the practitioner.

(e) A valid prescription or lawful order of a practitioner, in order to be effective in legalizing the possession of con-
trolled substances, must be issued in good faith for a legitimate medical purpose by one authorized to prescribe the use of such controlled substance. An order purporting to be a prescription not in the course of professional treatment is not a valid prescription or lawful order of a practitioner within the meaning and intent of this chapter; and the person who knows or should know that the person is filling such an order, as well as the person issuing it, can be charged with a violation of this chapter.

(f) A substance included in Schedule V must be distributed or dispensed only for a medical purpose.

(g) A practitioner may dispense or deliver a controlled substance to or for an individual or animal only for medical treatment or authorized research in the ordinary course of that practitioner’s profession. Medical treatment includes dispensing or administering a narcotic drug for pain, including intractable pain.

(h) No administrative sanction, or civil or criminal liability, authorized or created by this chapter may be imposed on a pharmacist for action taken in reliance on a reasonable belief that an order purporting to be a prescription was issued by a practitioner in the usual course of professional treatment or in authorized research.

(i) An individual practitioner may not dispense a substance included in Schedule II, III, or IV for that individual practitioner’s personal use. [2012 c 10 § 46; 2001 c 248 § 1; 1993 c 187 § 19; 1971 ex.s. c 308 § 69.50.308.]

Application—2012 c 10: See note following RCW 18.20.010.

69.50.309 Containers. A person to whom or for whose use any controlled substance has been prescribed, sold, or dispensed by a practitioner, and the owner of any animal for which such controlled substance has been prescribed, sold, or dispensed may lawfully possess it only in the container in which it was delivered to him or her by the person selling or dispensing the same. [2012 c 117 § 367; 1971 ex.s. c 308 § 69.50.309.]

69.50.310 Sodium pentobarbital—Registration of humane societies and animal control agencies for use in animal control. On and after September 21, 1977, a humane society and animal control agency may apply to the department for registration pursuant to the applicable provisions of this chapter for the sole purpose of being authorized to purchase, possess, and administer sodium pentobarbital to euthanize injured, sick, homeless, or unwanted domestic pets and animals. Any agency so registered shall not permit a person to administer sodium pentobarbital unless such person has demonstrated adequate knowledge of the potential hazards and proper techniques to be used in administering this drug. The department may issue a limited registration to carry out the provisions of this section. The board shall promulgate such rules as it deems necessary to insure strict compliance with the provisions of this section. The board may suspend or revoke registration upon determination that the person administering sodium pentobarbital has not demonstrated adequate knowledge as herein provided. This authority is granted in addition to any other power to suspend or revoke registration as provided by law. [1989 1st ex.s. c 9 § 435; 1977 ex.s. c 197 § 1.]

Additional notes found at www.leg.wa.gov

69.50.311 Triplicate prescription form program—Compliance by health care practitioners. Any licensed health care practitioner with prescription or dispensing authority shall, as a condition of licensure and as directed by the practitioner’s disciplinary board, consent to the requirement, if imposed, of complying with a triplicate prescription form program as may be established by rule by the department of health. [1989 1st ex.s. c 9 § 436; 1984 c 153 § 20.]

Additional notes found at www.leg.wa.gov

69.50.312 Electronic communication of prescription information—Board may adopt rules. (1) Information concerning an original prescription or information concerning a prescription refill for a controlled substance may be electronically communicated to a pharmacy of the patient’s choice pursuant to the provisions of this chapter if the electronically communicated prescription information complies with the following:

(a) Electronically communicated prescription information must comply with all applicable statutes and rules regarding the form, content, recordkeeping, and processing of a prescription for a legend drug;

(b) The system used for transmitting electronically communicated prescription information and the system used for receiving electronically communicated prescription information must be approved by the board. This subsection does not apply to currently used facsimile equipment transmitting an exact visual image of the prescription. The board shall maintain and provide, upon request, a list of systems used for electronically communicating prescription information currently approved by the board;

(c) An explicit opportunity for practitioners must be made to indicate their preference on whether a therapeutically equivalent generic drug may be substituted;

(d) Prescription drug orders are confidential health information, and may be released only to the patient or the patient’s authorized representative, the prescriber or other authorized practitioner then caring for the patient, or other persons specifically authorized by law to receive such information;

(e) To maintain confidentiality of prescription records, the electronic system shall have adequate security and systems safeguards designed to prevent and detect unauthorized access, modification, or manipulation of these records. The pharmacist in charge shall establish or verify the existence of policies and procedures which ensure the integrity and confidentiality of prescription information transmitted to the pharmacy by electronic means. All managers, employees, and agents of the pharmacy are required to read, sign, and comply with the established policies and procedures; and

(f) The pharmacist shall exercise professional judgment regarding the accuracy, validity, and authenticity of the prescription drug order received by way of electronic transmission, consistent with federal and state laws and rules and guidelines of the board.

(2) The board may adopt rules implementing this section. [1998 c 222 § 4.]

69.50.315 Medical assistance—Drug-related overdose—Naloxone—Prosecution for possession. (1)(a) A person acting in good faith who seeks medical assistance for
someone experiencing a drug-related overdose shall not be charged or prosecuted for possession of a controlled substance pursuant to RCW 69.50.4013, or penalized under RCW 69.50.4014, if the evidence for the charge of possession of a controlled substance was obtained as a result of the person seeking medical assistance.

(b) A person acting in good faith may receive a naloxone prescription, possess naloxone, and administer naloxone to an individual suffering from an apparent opiate-related overdose.

(2) A person who experiences a drug-related overdose and is in need of medical assistance shall not be charged or prosecuted for possession of a controlled substance pursuant to RCW 69.50.4013, or penalized under RCW 69.50.4014, if the evidence for the charge of possession of a controlled substance was obtained as a result of the overdose and the need for medical assistance.

(3) The protection in this section from prosecution for possession crimes under RCW 69.50.4013 shall not be grounds for suppression of evidence in other criminal charges. [2010 c 9 § 2.]

**Intent—2010 c 9:** "The legislature intends to save lives by increasing timely medical attention to drug overdose victims through the establishment of limited immunity from prosecution for people who seek medical assistance in a drug overdose situation. Drug overdose is the leading cause of unintentional injury death in Washington state, ahead of motor vehicle-related deaths. Washington state is one of sixteen states in which drug overdoses cause more deaths than traffic accidents. Drug overdose mortality rates have increased significantly since the 1990s, according to the centers for disease control and prevention, and illegal and prescription drug overdoses killed more than thirty-eight thousand people nationwide in 2006, the last year for which firm data is available. The Washington state department of health reports that in 1999 unintentional drug poisoning was responsible for four hundred three deaths in this state; in 2007, the number had increased to seven hundred sixty-one, compared with six hundred ten motor vehicle-related deaths that same year. Many drug overdose fatalities occur because peers delay or forego calling 911 for fear of arrest or police involvement, which researchers continually identify as the most significant barrier to the ideal first response of calling emergency services." [2010 c 9 § 1.]

### Article IV

**Offenses and Penalties**

69.50.401 **Prohibited acts: A—Penalties.** (1) Except as authorized by this chapter, it is unlawful for any person to manufacture, deliver, or possess with intent to manufacture or deliver, a controlled substance.

(2) Any person who violates this section with respect to:

(a) A controlled substance classified in Schedule I or II which is a narcotic drug or flunitrazepam, including its salts, isomers, and salts of isomers, classified in Schedule IV, is guilty of a class B felony and upon conviction may be imprisoned for not more than ten years, or (i) fined not more than twenty-five thousand dollars if the crime involved less than two kilograms of the drug, or both such imprisonment and fine; or (ii) if the crime involved two or more kilograms of the drug, then fined not more than one hundred thousand dollars for the first two kilograms and not more than fifty dollars for each gram in excess of two kilograms, or both such imprisonment and fine;

(b) Amphetamine, including its salts, isomers, and salts of isomers, or methamphetamine, including its salts, isomers, and salts of isomers, is guilty of a class B felony and upon conviction may be imprisoned for not more than ten years, or (i) fined not more than twenty-five thousand dollars if the crime involved less than two kilograms of the drug, or both such imprisonment and fine; or (ii) if the crime involved two or more kilograms of the drug, then fined not more than one hundred thousand dollars for the first two kilograms and not more than fifty dollars for each gram in excess of two kilograms, or both such imprisonment and fine.

(c) Any other controlled substance classified in Schedule I, II, or III, is guilty of a class C felony punishable according to chapter 9A.20 RCW;

(d) A substance classified in Schedule IV, except flunitrazepam, including its salts, isomers, and salts of isomers, is guilty of a class C felony punishable according to chapter 9A.20 RCW; or

(e) A substance classified in Schedule V, is guilty of a class C felony punishable according to chapter 9A.20 RCW.

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69.50.401 Counterfeit substances—Penalties. (1) Except as authorized by this chapter, it is unlawful for any person to create, deliver, or possess a counterfeit substance.

(2) Any person who violates this section with respect to:

(a) A counterfeit substance classified in Schedule I or II which is a narcotic drug, or flunitrazepam classified in Schedule IV, is guilty of a class B felony and upon conviction may be imprisoned for not more than ten years, fined not more than twenty-five thousand dollars, or both;

(b) A counterfeit substance which is methamphetamine, is guilty of a class B felony and upon conviction may be imprisoned for not more than ten years, fined not more than twenty-five thousand dollars, or both;

(c) Any other counterfeit substance classified in Schedule I, II, or III, is guilty of a class C felony punishable according to chapter 9A.20 RCW;

(d) A counterfeit substance classified in Schedule IV, except flunitrazepam, is guilty of a class C felony punishable according to chapter 9A.20 RCW;

(e) A counterfeit substance classified in Schedule V, is guilty of a class C felony punishable according to chapter 9A.20 RCW. [2003 c 53 § 332.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.50.4012 Delivery of substance in lieu of controlled substance—Penalty. (1) It is unlawful, except as authorized in this chapter and chapter 69.41 RCW, for any person to offer, arrange, or negotiate for the sale, gift, delivery, dispensing, distribution, or administration of a controlled substance to any person and then sell, give, deliver, dispense, distribute, or administer to that person any other liquid, substance, or material in lieu of such controlled substance.

(2) Any person who violates this section is guilty of a class C felony punishable according to chapter 9A.20 RCW. [2003 c 53 § 333.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.50.4013 Possession of controlled substance—Penalty. (1) It is unlawful for any person to possess a controlled substance unless the substance was obtained directly from, or pursuant to, a valid prescription or order of a practitioner while acting in the course of his or her professional practice, or except as otherwise authorized by this chapter.

(2) Except as provided in RCW 69.50.4014, any person who violates this section is guilty of a class C felony punishable under chapter 9A.20 RCW. [2003 c 53 § 334.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.50.4014 Possession of forty grams or less of marihuana—Penalty. Except as provided in RCW 69.50.401(2)(c), any person found guilty of possession of forty grams or less of marihuana is guilty of a misdemeanor. [2003 c 53 § 335.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.50.4015 Involving a person under eighteen in unlawful controlled substance transaction—Penalty. (1) It is unlawful to compensate, threaten, solicit, or in any other manner involve a person under the age of eighteen years in a transaction unlawfully to manufacture, sell, or deliver a controlled substance.

(2) A violation of this section is a class C felony punishable according to chapter 9A.20 RCW. [2003 c 53 § 336.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.50.4016 Provisions not applicable to offenses under RCW 69.50.410. RCW 69.50.401 through 69.50.405 shall not apply to offenses defined and punishable under the provisions of RCW 69.50.410. [2003 c 53 § 337.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.50.402 Prohibited acts: B—Penalties. (1) It is unlawful for any person:

(a) Who is subject to Article III to distribute or dispense a controlled substance in violation of RCW 69.50.308;

(b) Who is a registrant, to manufacture a controlled substance not authorized by his or her registration, or to distribute or dispense a controlled substance not authorized by his or her registration to another registrant or other authorized person;

(c) Who is a practitioner, to prescribe, order, dispense, administer, supply, or give to any person:

(i) Any amphetamine, including its salts, optical isomers, and salts of optical isomers classified as a schedule II controlled substance by the board of pharmacy pursuant to chapter 34.05 RCW; or

(ii) Any nonnarcotic stimulant classified as a schedule II controlled substance and designated as a nonnarcotic stimulant by the board of pharmacy pursuant to chapter 34.05 RCW; except for the treatment of narcolepsy or for the treatment of hyperkinesis, or for the treatment of drug-induced brain dysfunction, or for the treatment of epilepsy, or for the differential diagnostic psychiatric evaluation of depression, or for the treatment of depression shown to be refractory to other therapeutic modalities, or for the treatment of multiple sclerosis, or for the clinical investigation of the effects of such drugs or compounds, in which case an investigational protocol therefor shall have been submitted to and reviewed and approved by the state board of pharmacy before the investigation has been begun: PROVIDED, That the board of pharmacy, in consultation with the medical quality assurance commission and the osteopathic disciplinary board, may establish by rule, pursuant to chapter 34.05 RCW, disease states or conditions in addition to those listed in this subsection for the treatment of which Schedule II nonnarcotic stimulants may be prescribed, ordered, dispensed, administered, supplied, or given to patients by practitioners: AND PROVIDED, FURTHER, That investigations by the board of pharmacy of abuse of prescriptive authority by physicians, licensed pursuant to chapter 18.71 RCW, pursuant to subsection (1)(c) of this section
shall be done in consultation with the medical quality assurance commission;

(d) To refuse or fail to make, keep or furnish any record, notification, order form, statement, invoice, or information required under this chapter;

(e) To refuse an entry into any premises for any inspection authorized by this chapter; or

(f) Knowingly to keep or maintain any store, shop, warehouse, dwelling, building, vehicle, boat, aircraft, or other structure or place, which is resorted to by persons using controlled substances in violation of this chapter for the purpose of using these substances, or which is used for keeping or selling them in violation of this chapter.

(2) Any person who violates this section is guilty of a class C felony and upon conviction may be imprisoned for not more than two years, fined not more than two thousand dollars, or both. [2010 c 177 § 7; 2003 c 53 § 338; 1994 sp.s. c 9 § 740; 1980 c 138 § 6; 1979 ex.s. c 119 § 1; 1971 ex.s. c 308 § 69.50.402.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

Additional notes found at www.leg.wa.gov

69.50.403 Prohibited acts: C—Penalties. (1) It is unlawful for any person knowingly or intentionally:

(a) To distribute as a registrant a controlled substance classified in Schedules I or II, except pursuant to an order form as required by *RCW 69.50.307;

(b) To use in the course of the manufacture, distribution, or dispensing of a controlled substance, or to use for the purpose of acquiring or obtaining a controlled substance, a registration number which is fictitious, revoked, suspended, or issued to another person;

(c) To obtain or attempt to obtain a controlled substance, or procure or attempt to procure the administration of a controlled substance, (i) by fraud, deceit, misrepresentation, or subterfuge; or (ii) by forgery or alteration of a prescription or any written order; or (iii) by the concealment of material fact; or (iv) by the use of a false name or the giving of a false address;

(d) To falsely assume the title of, or represent herself or himself to be, a manufacturer, wholesaler, pharmacist, physician, dentist, veterinarian, or other authorized person for the purpose of obtaining a controlled substance;

(e) To make or utter any false or forged prescription or false or forged written order;

(f) To affix any false or forged label to a package or receptacle containing controlled substances;

(g) To furnish false or fraudulent material information in, or omit any material information from, any application, report, or other document required to be kept or filed under this chapter, or any record required to be kept by this chapter;

(h) To possess a false or fraudulent prescription with intent to obtain a controlled substance; or

(i) To attempt to illegally obtain controlled substances by providing more than one name to a practitioner when obtaining a prescription for a controlled substance. If a person’s name is legally changed during the time period that he or she is receiving health care from a practitioner, the person shall inform all providers of care so that the medical and pharmacy records for the person may be filed under a single name identifier.

(2) Information communicated to a practitioner in an effort unlawfully to procure a controlled substance or unlawfully to procure the administration of such substance, shall not be deemed a privileged communication.

(3) A person who violates this section is guilty of a class C felony and upon conviction may be imprisoned for not more than two years, or fined not more than two thousand dollars, or both. [2003 c 53 § 339; 1996 c 255 § 1; 1993 c 187 § 21; 1971 ex.s. c 308 § 69.50.403.]

*Reviser’s note: RCW 69.50.307 was repealed by 2001 c 248 § 2.

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.50.404 Penalties under other laws. Any penalty imposed for violation of this chapter is in addition to, and not in lieu of, any civil or administrative penalty or sanction otherwise authorized by law. [1971 ex.s. c 308 § 69.50.404.]

69.50.405 Bar to prosecution. If a violation of this chapter is a violation of a federal law or the law of another state, a conviction or acquittal under federal law or the law of another state for the same act is a bar to prosecution in this state. [1971 ex.s. c 308 § 69.50.405.]

69.50.406 Distribution to persons under age eighteen. (1) Any person eighteen years of age or over who violates RCW 69.50.401 by distributing a controlled substance listed in Schedules I or II which is a narcotic drug or methamphetamine, including its salts, isomers, and salts of isomers, or flunitrazepam, including its salts, isomers, and salts of isomers, listed in Schedule IV, to a person under eighteen years of age is guilty of a class A felony punishable by the fine authorized by RCW 69.50.401(2)(a) or (b), by a term of imprisonment up to twice that authorized by RCW 69.50.401(2)(a) or (b), or by both.

(2) Any person eighteen years of age or over who violates RCW 69.50.401 by distributing any other controlled substance listed in Schedules I, II, III, IV, and V to a person under eighteen years of age who is at least three years his or her junior is guilty of a class B felony punishable by the fine authorized by RCW 69.50.401(2)(c), (d), or (e), by a term of imprisonment up to twice that authorized by RCW 69.50.401(2)(c), (d), or (e), or both. [2005 c 218 § 2; 2003 c 53 § 340; 1998 c 290 § 2; 1996 c 205 § 7; 1987 c 458 § 5; 1971 ex.s. c 308 § 69.50.406.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

Additional notes found at www.leg.wa.gov

69.50.407 Conspiracy. Any person who attempts or conspires to commit any offense defined in this chapter is punishable by imprisonment or fine or both which may not exceed the maximum punishment prescribed for the offense, the commission of which was the object of the attempt or conspiracy. [1971 ex.s. c 308 § 69.50.407.]

69.50.408 Second or subsequent offenses. (1) Any person convicted of a second or subsequent offense under this chapter may be imprisoned for a term up to twice the
term otherwise authorized, fined an amount up to twice that otherwise authorized, or both.

(2) For purposes of this section, an offense is considered a second or subsequent offense, if, prior to his or her conviction of the offense, the offender has at any time been convicted under this chapter or under any statute of the United States or of any state relating to narcotic drugs, marihuana, depressant, stimulant, or hallucinogenic drugs.

(3) This section does not apply to offenses under RCW 69.50.413. [2003 c 53 § 341; 1989 c 8 § 3; 1971 ex.s. c 308 § 69.50.408.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.50.410 Prohibited acts: D—Penalties. (1) Except as authorized by this chapter it is a class C felony for any person to sell for profit any controlled substance or counterfeit substance classified in Schedule I, RCW 69.50.204, except leaves and flowering tops of marihuana.

For the purposes of this section only, the following words and phrases shall have the following meanings:

(a) "To sell" means the passing of title and possession of a controlled substance from the seller to the buyer for a price whether or not the price is paid immediately or at a future date.

(b) "For profit" means the obtaining of anything of value in exchange for a controlled substance.

(c) "Price" means anything of value.

(2) (a) Any person convicted of a violation of subsection (1) of this section shall receive a sentence of not more than five years in a correctional facility of the department of social and health services for the first offense.

(b) Any person convicted on a second or subsequent cause, the sale having transpired after prosecution and conviction on the first cause, of subsection (1) of this section shall receive a mandatory sentence of five years in a correctional facility of the department of social and health services and no judge of any court shall suspend or defer the sentence imposed for the second or subsequent violation of subsection (1) of this section.

(3) (a) Any person convicted of a violation of subsection (1) of this section by selling heroin shall receive a mandatory sentence of two years in a correctional facility of the department of social and health services and no judge of any court shall suspend or defer the sentence imposed for such violation.

(b) Any person convicted on a second or subsequent sale of heroin, the sale having transpired after prosecution and conviction on the first cause of the sale of heroin shall receive a mandatory sentence of ten years in a correctional facility of the department of social and health services and no judge of any court shall suspend or defer the sentence imposed for this second or subsequent violation: PROVIDED, That the indeterminate sentence review board under RCW 9.95.040 shall not reduce the minimum term imposed for a violation under this subsection.

(4) Whether or not a mandatory minimum term has expired, an offender serving a sentence under this section may be granted an extraordinary medical placement when authorized under *RCW 9.94A.728(4).

(5) In addition to the sentences provided in subsection (2) of this section, any person convicted of a violation of subsection (1) of this section shall be fined in an amount calculated to at least eliminate any and all proceeds or profits directly or indirectly gained by such person as a result of sales of controlled substances in violation of the laws of this or other states, or the United States, up to the amount of five hundred thousand dollars on each count.

(6) Any person, addicted to the use of controlled substances, who voluntarily applies to the department of social and health services for the purpose of participating in a rehabilitation program approved by the department for addicts of controlled substances shall be immune from prosecution for subsection (1) offenses unless a filing of an information or indictment against such person for a violation of subsection (1) of this section is made prior to his or her voluntary participation in the program of the department of social and health services. All applications for immunity under this section shall be sent to the department of social and health services in Olympia. It shall be the duty of the department to stamp each application received pursuant to this section with the date and time of receipt.

(7) This section shall not apply to offenses defined and punishable under the provisions of RCW 69.50.401 through 69.50.4015. [2003 c 53 § 342; 1999 c 324 § 6; 1975-'76 2nd ex.s. c 103 § 1; 1973 2nd ex.s. c 2 § 2.]

*Reviser's note: RCW 9.94A.728 was amended by 2009 c 455 § 2, changing subsection (4) to subsection (3).

69.50.412 Prohibited acts: E—Penalties. (1) It is unlawful for any person to use drug paraphernalia to plant, propagate, cultivate, grow, harvest, manufacture, compound, convert, produce, process, prepare, test, analyze, pack, repack, store, contain, conceal, inject, ingest, inhale, or otherwise introduce into the human body a controlled substance. Any person who violates this subsection is guilty of a misdemeanor.

(2) It is unlawful for any person to deliver, possess with intent to deliver, or manufacture with intent to deliver drug paraphernalia, knowing, or under circumstances where one reasonably should know, that it will be used to plant, propagate, cultivate, grow, harvest, manufacture, compound, convert, produce, process, prepare, test, analyze, pack, repack, store, contain, conceal, inject, ingest, inhale, or otherwise introduce into the human body a controlled substance. Any person who violates this subsection is guilty of a misdemeanor.

(3) Any person eighteen years of age or over who violates subsection (2) of this section by delivering drug paraphernalia to a person under eighteen years of age who is at least three years his or her junior is guilty of a gross misdemeanor.

(4) It is unlawful for any person to place in any newspaper, magazine, handbill, or other publication any advertisement, knowing, or under circumstances where one reasonably should know, that the purpose of the advertisement, in whole or in part, is to promote the sale of objects designed or intended for use as drug paraphernalia. Any person who violates this subsection is guilty of a misdemeanor.

2012 [Title 69 RCW—page 85]
(5) It is lawful for any person over the age of eighteen to possess sterile hypodermic syringes and needles for the purpose of reducing bloodborne diseases. [2012 c 117 § 368; 2002 c 213 § 1; 1981 c 48 § 2.]

Additional notes found at www.leg.wa.gov

69.50.4121 Drug paraphernalia—Selling or giving—Penalty. (1) Every person who sells or gives, or permits to be sold or given to any person any drug paraphernalia in any form commits a class 1 civil infraction under chapter 7.80 RCW. For purposes of this subsection, "drug paraphernalia" means all equipment, products, and materials of any kind which are used, intended for use, or designed for use in planting, propagating, cultivating, growing, harvesting, manufacturing, compounding, converting, producing, processing, preparing, testing, analyzing, packaging, repackaging, storing, containing, concealing, injecting, ingesting, inhaling, or otherwise introducing into the human body a controlled substance. Drug paraphernalia includes, but is not limited to objects used, intended for use, or designed for use in ingesting, inhaling, or otherwise introducing marijuana, cocaine, hashish, or hashish oil into the human body, such as:

(a) Metal, wooden, acrylic, glass, stone, plastic, or ceramic pipes with or without screens, permanent screens, hashish heads, or punctured metal bowls;
(b) Water pipes;
(c) Carburetion tubes and devices;
(d) Smoking and carburetion masks;
(e) Roach clips: Meaning objects used to hold burning material, such as a marijuana cigarette, that has become too small or too short to be held in the hand;
(f) Miniature cocaine spoons and cocaine vials;
(g) Chamber pipes;
(h) Carburetor pipes;
(i) Electric pipes;
(j) Air-driven pipes;
(k) Chillums;
(l) Bongs; and
(m) Ice pipes or chillers.
(2) It shall be no defense to a prosecution for a violation of this section that the person acted, or was believed by the defendant to act, as agent or representative of another.
(3) Nothing in subsection (1) of this section prohibits legal distribution of injection syringe equipment through public health and community based HIV prevention programs, and pharmacies. [2002 c 213 § 2; 1998 c 317 § 1.]

69.50.413 Health care practitioners—Suspension of license for violation of chapter. The license of any licensed health care practitioner shall be suspended for any violation of this chapter. The suspension shall run concurrently with, and not less than, the term of the sentence for the violation. [1984 c 153 § 21.]

69.50.414 Sale or transfer of controlled substance to minor—Cause of action by parent—Damages. The parent or legal guardian of any minor to whom a controlled substance, as defined in RCW 69.50.101, is sold or transferred, shall have a cause of action against the person who sold or transferred the controlled substance for all damages to the minor or his or her parent or legal guardian caused by such

sale or transfer. Damages shall include: (a) Actual damages, including the cost for treatment or rehabilitation of the minor child's drug dependency, (b) forfeiture to the parent or legal guardian of the cash value of any proceeds received from such sale or transfer of a controlled substance, and (e) reasonable attorney fees.

This section shall not apply to a practitioner, as defined in *RCW 69.50.101(t), who sells or transfers a controlled substance to a minor pursuant to a valid prescription or order. [1986 c 124 § 10.]

*Reviser's note: The reference to RCW 69.50.101(t) is erroneous. Practitioner" is defined in (w) of that section. RCW 69.50.101 was alphabetized pursuant to RCW 1.08.015(2)(k), changing subsection (w) to subsection (x).

69.50.415 Controlled substances homicide—Penalty. (1) A person who unlawfully delivers a controlled substance in violation of RCW 69.50.401(2) (a), (b), or (c) which controlled substance is subsequently used by the person to whom it was delivered, resulting in the death of the user, is guilty of controlled substances homicide.
(2) Controlled substances homicide is a class B felony punishable according to chapter 9A.20 RCW. [2003 c 53 § 343; 1996 c 205 § 8; 1987 c 458 § 2.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

Additional notes found at www.leg.wa.gov

69.50.416 Counterfeit substances prohibited—Penalties. (1) It is unlawful for any person knowingly or intentionally to manufacture, deliver, or possess with intent to manufacture or deliver, a controlled substance which, or the container or labeling of which, without authorization, bears the trademark, trade name, or other identifying mark, imprint, number, or device, or any likeness thereof, of a manufacturer, distributor, or dispenser, other than the person who in fact manufactured, distributed, or dispensed the substance.
(2) It is unlawful for any person knowingly or intentionally to make, distribute, or possess a punch, die, plate, stone, or other thing designed to print, imprint, or reproduce the trademark, trade name, or other identifying mark, imprint, or device of another or any likeness of any of the foregoing upon any drug or container or labeling thereof.
(3) A person who violates this section is guilty of a class C felony and upon conviction may be imprisoned for not more than two years, fined not more than two thousand dollars, or both. [2003 c 53 § 344; 1993 c 187 § 22.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.50.420 Violations—Juvenile driving privileges.
(1) If a juvenile thirteen years of age or older and under the age of twenty-one is found by a court to have committed any offense that is a violation of this chapter, the court shall notify the department of licensing within twenty-four hours after entry of the judgment.
(2) Except as otherwise provided in subsection (3) of this section, upon petition of a juvenile whose privilege to drive has been revoked pursuant to RCW 46.20.265, the court may at any time the court deems appropriate notify the department of licensing to reinstate the juvenile’s privilege to drive.
(3) If the conviction is for the juvenile’s first violation of this chapter or chapter 66.44, 69.41, or 69.52 RCW, the juvenile may not petition the court for reinstatement of the juvenile’s privilege to drive revoked pursuant to RCW 46.20.265 until the later of ninety days after the date the juvenile turns sixteen or ninety days after the judgment was entered. If the conviction was for the juvenile’s second or subsequent violation of this chapter or chapter 66.44, 69.41, or 69.52 RCW, the juvenile may not petition the court for reinstatement of the juvenile’s privilege to drive revoked pursuant to RCW 46.20.265 until the later of the date the juvenile turns seventeen or one year after the date judgment was entered. [1989 c 271 § 120; 1988 c 148 § 5.]

Legislative finding—Severability—1988 c 148: See notes following RCW 13.40.265.

Additional notes found at www.leg.wa.gov

69.50.425 Misdemeanor violations—Minimum penalties. A person who is convicted of a misdemeanor violation of any provision of this chapter shall be punished by imprisonment for not less than twenty-four consecutive hours, and by a fine of not less than two hundred fifty dollars. On a second or subsequent conviction, the fine shall not be less than five hundred dollars. These fines shall be in addition to any other fine or penalty imposed. Unless the court finds that the imposition of the minimum imprisonment will pose a substantial risk to the defendant’s physical or mental well-being or that local jail facilities are in an overcrowded condition, the minimum term of imprisonment shall not be suspended or deferred. If the court finds such risk or overcrowding exists, it shall sentence the defendant to a minimum of forty hours of community restitution. If a minimum term of imprisonment is suspended or deferred, the court shall state in writing the reason for granting the suspension or deferral and the facts upon which the suspension or deferral is based. Unless the court finds the person to be indigent, the minimum fine shall not be suspended or deferred. [2002 c 175 § 44; 1989 c 271 § 105.]

Effective date—2002 c 175: See note following RCW 7.80.130.

Additional notes found at www.leg.wa.gov

69.50.430 Additional fine for certain felony violations. (1) Every person convicted of a felony violation of RCW 69.50.401 through 69.50.4013, 69.50.4015, 69.50.402, 69.50.403, 69.50.406, 69.50.407, 69.50.410, or 69.50.415 shall be fined one thousand dollars in addition to any other fine or penalty imposed. Unless the court finds the person to be indigent, this additional fine shall not be suspended or deferred by the court.

(2) On a second or subsequent conviction for violation of any of the laws listed in subsection (1) of this section, the person shall be fined two thousand dollars in addition to any other fine or penalty imposed. Unless the court finds the person to be indigent, this additional fine shall not be suspended or deferred by the court. [2003 c 53 § 345; 1989 c 271 § 106.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

Additional notes found at www.leg.wa.gov

69.50.435 Violations committed in or on certain public places or facilities—Additional penalty—Defenses—

Construction—Definitions. (1) Any person who violates RCW 69.50.401 by manufacturing, selling, delivering, or possessing with the intent to manufacture, sell, or deliver a controlled substance listed under RCW 69.50.401 or who violates RCW 69.50.410 by selling for profit any controlled substance or counterfeit substance classified in schedule I, RCW 69.50.204, except leaves and flowering tops of marijuana to a person:

(a) In a school;
(b) On a school bus;
(c) Within one thousand feet of a school bus route stop designated by the school district;
(d) Within one thousand feet of the perimeter of the school grounds;
(e) In a public park;
(f) In a public housing project designated by a local governing authority as a drug-free zone;
(g) On a public transit vehicle;
(h) In a public transit stop shelter;
(i) At a civic center designated as a drug-free zone by the local governing authority; or
(j) Within one thousand feet of the perimeter of a facility designated under (i) of this subsection, if the local governing authority specifically designates the one thousand foot perimeter may be punished by a fine of up to twice the fine otherwise authorized by this chapter, but not including twice the fine authorized by RCW 69.50.406, or by imprisonment of up to twice the imprisonment otherwise authorized by this chapter, but not including twice the imprisonment authorized by RCW 69.50.406, or by both such fine and imprisonment. The provisions of this section shall not operate to more than double the fine or imprisonment otherwise authorized by this chapter for an offense.

(2) It is not a defense to a prosecution for a violation of this section that the person was unaware that the prohibited conduct took place while in a school or school bus or within one thousand feet of the school or school bus route stop, in a public park, in a public housing project designated by a local governing authority as a drug-free zone, on a public transit vehicle, in a public transit stop shelter, at a civic center designated as a drug-free zone by the local governing authority, or within one thousand feet of the perimeter of a facility designated under subsection (1)(i) of this section, if the local governing authority specifically designates the one thousand foot perimeter.

(3) It is not a defense to a prosecution for a violation of this section or any other prosecution under this chapter that persons under the age of eighteen were not present in the school, the school bus, the public park, the public housing project designated by a local governing authority as a drug-free zone, or the public transit vehicle, or at the school bus route stop, the public transit vehicle stop shelter, at a civic center designated as a drug-free zone by the local governing authority, or within one thousand feet of the perimeter of a facility designated under subsection (1)(i) of this section, if the local governing authority specifically designates the one thousand foot perimeter at the time of the offense or that school was not in session.

(4) It is an affirmative defense to a prosecution for a violation of this section that the prohibited conduct took place
entirely within a private residence, that no person under eighteen years of age or younger was present in such private residence at any time during the commission of the offense, and that the prohibited conduct did not involve delivering, manufacturing, selling, or possessing with the intent to manufacture, sell, or deliver any controlled substance in RCW 69.50.401 for profit. The affirmative defense established in this section shall be proved by the defendant by a preponderance of the evidence. This section shall not be construed to establish an affirmative defense with respect to a prosecution for an offense defined in any other section of this chapter.

(5) In a prosecution under this section, a map produced or reproduced by any municipality, school district, county, transit authority engineer, or public housing authority for the purpose of depicting the location and boundaries of the area on or within one thousand feet of any property used for a school, school bus route stop, public park, public housing project designated by a local governing authority as a drug-free zone, public transit vehicle stop shelter, or a civic center designated as a drug-free zone by a local governing authority, or a true copy of such a map, shall under proper authentication, be admissible and shall constitute prima facie evidence of the location and boundaries of those areas if the governing body of the municipality, school district, county, or transit authority has adopted a resolution or ordinance approving the map as the official location and record of the location and boundaries of the area on or within one thousand feet of the school, school bus route stop, public park, public housing project designated by a local governing authority as a drug-free zone, public transit vehicle stop shelter, or civic center designated as a drug-free zone by a local governing authority. Any map approved under this section or a true copy of the map shall be filed with the clerk of the municipality or county, and shall be maintained as an official record of the municipality or county. This section shall not be construed as precluding the prosecution from introducing or relying upon any other evidence or testimony to establish any element of the offense. This section shall not be construed as precluding the use or admissibility of any map or diagram other than the one which has been approved by the governing body of a municipality, school district, county, transit authority, or public housing authority if the map or diagram is otherwise admissible under court rule.

(6) As used in this section the following terms have the meanings indicated unless the context clearly requires otherwise:

(a) "School" has the meaning under RCW 28A.150.010 or 28A.150.020. The term "school" also includes a private school approved under RCW 28A.195.010;

(b) "School bus" means a school bus as defined by the superintendent of public instruction by rule which is owned and operated by any school district and all school buses which are privately owned and operated under contract or otherwise with any school district in the state for the transportation of students. The term does not include buses operated by common carriers in the urban transportation of students such as transportation of students through a municipal transportation system;

(c) "School bus route stop" means a school bus stop as designated by a school district;

(d) "Public park" means land, including any facilities or improvements on the land, that is operated as a park by the state or a local government;

(e) "Public transit vehicle" means any motor vehicle, streetcar, train, trolley vehicle, or any other device, vessel, or vehicle which is owned or operated by a transit authority and which is used for the purpose of carrying passengers on a regular schedule;

(f) "Transit authority" means a city, county, or state transportation system, transportation authority, public transportation benefit area, public transit authority, or metropolitan municipal corporation within the state that operates public transit vehicles;

(g) "Stop shelter" means a passenger shelter designated by a transit authority;

(h) "Civic center" means a publicly owned or publicly operated place or facility used for recreational, educational, or cultural activities;

(i) "Public housing project" means the same as "housing project" as defined in RCW 35.82.020. [2003 c 53 § 346. Prior: 1997 c 30 § 2; 1997 c 23 § 1; 1996 c 14 § 2; 1991 c 32 § 4; prior: 1990 c 244 § 1; 1990 c 33 § 588; 1989 c 271 § 112.]

Purpose—Statutory references—Severability—1990 c 244 § 1: See notes following RCW 2.48.180.

Findings—Intent—1997 c 30: "The legislature finds that a large number of illegal drug transactions occur in or near public housing projects. The legislature also finds that this activity places the families and children residing in these housing projects at risk for drug-related crimes and increases the general level of fear among the residents of the housing project and the areas surrounding these projects. The intent of the legislature is to allow local governments to designate public housing projects as drug-free zones." [1997 c 30 § 1.]

Findings—Intent—1996 c 14: "The legislature finds that a large number of illegal drug transactions occur in or near publicly owned places used for recreational, educational, and cultural purposes. The legislature also finds that this activity places the people using these facilities at risk for drug-related crimes, discourages the use of recreational, educational, and cultural facilities, blights the economic development around these facilities, and increases the general level of fear among the residents of the areas surrounding these facilities. The intent of the legislature is to allow local governments to designate a perimeter of one thousand feet around publicly owned places used primarily for recreation, education, and cultural activities as drug-free zones." [1996 c 14 § 1.]

Penalty. (1) It is unlawful for any person to possess ephedrine or any of its salts or isomers or salts of isomers, pseudoephedrine or any of its salts or isomers or salts of isomers, pressurized ammonia gas, or pressurized ammonia gas solution with intent to manufacture methamphetamine, including its salts, isomers, and salts of isomers.

(2) Any person who violates this section is guilty of a class B felony and may be imprisoned for not more than ten years, fined not more than twenty-five thousand dollars, or both. Three thousand dollars of the fine may not be suspended. As collected, the first three thousand dollars of the fine must be deposited with the law enforcement agency having responsibility for cleanup of laboratories, sites, or substances used in the manufacture of the methamphetamine, including its salts, isomers, and salts of isomers. The fine
moneys deposited with that law enforcement agency must be used for such clean-up cost. [2005 c 218 § 3; 2003 c 53 § 347; 2002 c 134 § 1; 2000 c 225 § 4; 1997 c 71 § 3; 1996 c 205 § 1.]

**Intent—Effective date—2003 c 53:** See notes following RCW 2.28.180.

**Effective date—2002 c 134:** "This act is necessary for the immediate preservation of the public peace, health, or safety, or support of the state government and its existing public institutions, and takes effect immediately [March 26, 2002]." [2002 c 134 § 5.]

Additional notes found at www.leg.wa.gov

### ARTICLE V

#### ENFORCEMENT AND ADMINISTRATIVE PROVISIONS

**69.50.500 Powers of enforcement personnel.** (a) It is hereby made the duty of the state board of pharmacy, the department, and their officers, agents, inspectors and representatives, and all law enforcement officers within the state, and of all prosecuting attorneys, to enforce all provisions of this chapter, except those specifically delegated, and to cooperate with all agencies charged with the enforcement of the laws of the United States, of this state, and all other states, relating to controlled substances as defined in this chapter.

(b) Employees of the department of health, who are so designated by the board as enforcement officers are declared to be peace officers and shall be vested with police powers to enforce the drug laws of this state, including this chapter. [1989 1st ex.s. c 9 § 437; 1971 ex.s. c 308 § 69.50.000.]

Additional notes found at www.leg.wa.gov

**69.50.501 Administrative inspections.** The state board of pharmacy may make administrative inspections of controlled premises in accordance with the following provisions:

1. For purposes of this section only, "controlled premises" means:
   (a) places where persons registered or exempted from registration requirements under this chapter are required to keep records; and
   (b) places including factories, warehouses, establishments, and conveyances in which persons registered or exempted from registration requirements under this chapter are permitted to hold, manufacture, compound, process, sell, deliver, or otherwise dispose of any controlled substance.

2. When authorized by an administrative inspection warrant issued pursuant to RCW 69.50.502 an officer or employee designated by the board, upon presenting the warrant and appropriate credentials to the owner, operator, or agent in charge, may enter controlled premises for the purpose of conducting an administrative inspection.

3. When authorized by an administrative inspection warrant, an officer or employee designated by the board may:
   (a) inspect and copy records required by this chapter to be kept;
   (b) inspect, within reasonable limits and in a reasonable manner, controlled premises and all pertinent equipment, finished and unfinished material, containers and labeling found therein, and, except as provided in subsection (5) of this section, all other things therein, including records, files, papers, processes, controls, and facilities bearing on violation of this chapter; and
   (c) inventory any stock of any controlled substance therein and obtain samples thereof;

4. This section does not prevent the inspection without a warrant of books and records pursuant to an administrative subpoena issued in accordance with chapter 34.05 RCW, nor does it prevent entries and administrative inspections, including seizures of property, without a warrant:
   (a) if the owner, operator, or agent in charge of the controlled premises consents;
   (b) in situations presenting imminent danger to health or safety;
   (c) in situations involving inspection of conveyances if there is reasonable cause to believe that the mobility of the conveyance makes it impracticable to obtain a warrant;
   (d) in any other exceptional or emergency circumstance where time or opportunity to apply for a warrant is lacking; or,
   (e) in all other situations in which a warrant is not constitutionally required;

5. An inspection authorized by this section shall not extend to financial data, sales data, other than shipment data, or pricing data unless the owner, operator, or agent in charge of the controlled premises consents in writing. [1971 ex.s. c 308 § 69.50.01.]

**69.50.502 Warrants for administrative inspections.** Issuance and execution of administrative inspection warrants shall be as follows:

1. A judge of a superior court, or a judge of a district court within his or her jurisdiction, and upon proper oath or affirmation showing probable cause, may issue warrants for the purpose of conducting administrative inspections authorized by this chapter or rules hereunder, and seizures of property appropriate to the inspections. For purposes of the issuance of administrative inspection warrants, probable cause exists upon showing a valid public interest in the effective enforcement of this chapter or rules hereunder, sufficient to justify administrative inspection of the area, premises, building, or conveyance in the circumstances specified in the application for the warrant;

2. A warrant shall issue only upon an affidavit of a designated officer or employee having knowledge of the facts alleged, sworn to before the judge and establishing the grounds for issuing the warrant. If the judge is satisfied that grounds for the application exist or that there is probable cause to believe they exist, he or she shall issue a warrant identifying the area, premises, building, or conveyance to be inspected, the purpose of the inspection, and, if appropriate, the type of property to be inspected, if any. The warrant shall:
   (a) State the grounds for its issuance and the name of each person whose affidavit has been taken in support thereof;
   (b) Be directed to a person authorized by RCW 69.50.500 to execute it;
   (c) Command the person to whom it is directed to inspect the area, premises, building, or conveyance identified for the purpose specified and, if appropriate, direct the seizure of the property specified;
(d) Identify the item or types of property to be seized, if any;
(e) Direct that it be served during normal business hours and designate the judge to whom it shall be returned;
(3) A warrant issued pursuant to this section must be executed and returned within ten days of its date unless, upon a showing of a need for additional time, the court orders otherwise. If property is seized pursuant to a warrant, a copy shall be given to the person from whom or from whose premises the property is taken, together with a receipt for the property taken. The return of the warrant shall be made promptly, accompanied by a written inventory of any property taken. The inventory shall be made in the presence of the person executing the warrant and of the person from whose possession or premises the property was taken, if present, or in the presence of at least one credible person other than the person executing the warrant. A copy of the inventory shall be delivered to the person from whom or from whose premises the property was taken and to the applicant for the warrant;
(4) The judge who has issued a warrant shall attach thereto a copy of the return and all papers returnable in connection therewith and file them with the clerk of the court in which the inspection was made. [2012 c 117 § 369; 1971 ex.s. c 308 § 69.50.502.]

69.50.503 Injunctions. (a) The superior courts of this state have jurisdiction to restrain or enjoin violations of this chapter.
(b) The defendant may demand trial by jury for an alleged violation of an injunction or restraining order under this section. [1971 ex.s. c 308 § 69.50.503.]

69.50.504 Cooperative arrangements. The state board of pharmacy shall cooperate with federal and other state agencies in discharging its responsibilities concerning traffic in controlled substances and in suppressing the abuse of controlled substances. [1971 ex.s. c 308 § 69.50.504.]

69.50.505 Seizure and forfeiture. (1) The following are subject to seizure and forfeiture and no property right exists in them:
(a) All controlled substances which have been manufactured, distributed, dispensed, acquired, or possessed in violation of this chapter or chapter 69.41 or 69.52 RCW, and all hazardous chemicals, as defined in RCW 64.44.010, used or intended to be used in the manufacture of controlled substances;
(b) All raw materials, products, and equipment of any kind which are used, or intended for use, in manufacturing, compounding, processing, delivering, importing, or exporting any controlled substance in violation of this chapter or chapter 69.41 or 69.52 RCW;
(c) All property which is used, or intended for use, as a container for property described in (a) or (b) of this subsection;
(d) All conveyances, including aircraft, vehicles, or vessels, which are used, or intended for use, in any manner to facilitate the sale, delivery, or receipt of property described in (a) or (b) of this subsection, except that:
   (i) No conveyance used by any person as a common carrier in the transaction of business as a common carrier is subject to forfeiture under this section unless it appears that the owner or other person in charge of the conveyance is a consenting party or privy to a violation of this chapter or chapter 69.41 or 69.52 RCW;
   (ii) No conveyance is subject to forfeiture under this section by reason of any act or omission established by the owner thereof to have been committed or omitted without the owner’s knowledge or consent;
   (iii) No conveyance is subject to forfeiture under this section if used in the receipt of only an amount of marijuana for which possession constitutes a misdemeanor under RCW 69.50.4014;
   (iv) A forfeiture of a conveyance encumbered by a bona fide security interest is subject to the interest of the secured party if the secured party neither had knowledge of nor consented to the act or omission; and
   (v) When the owner of a conveyance has been arrested under this chapter or chapter 69.41 or 69.52 RCW the conveyance in which the person is arrested may not be subject to forfeiture unless it is seized or process is issued for its seizure within ten days of the owner’s arrest;
   (e) All books, records, and research products and materials, including formulas, microfilm, tapes, and data which are used, or intended for use, in violation of this chapter or chapter 69.41 or 69.52 RCW;
   (f) All drug paraphernalia;
   (g) All moneys, negotiable instruments, securities, or other tangible or intangible property of value furnished or intended to be furnished by any person in exchange for a controlled substance in violation of this chapter or chapter 69.41 or 69.52 RCW, all tangible or intangible personal property, proceeds, or assets acquired in whole or in part with proceeds traceable to an exchange or series of exchanges in violation of this chapter or chapter 69.41 or 69.52 RCW, and all moneys, negotiable instruments, and securities used or intended to be used to facilitate any violation of this chapter or chapter 69.41 or 69.52 RCW. A forfeiture of money, negotiable instruments, securities, or other tangible or intangible property encumbered by a bona fide security interest is subject to the interest of the secured party if, at the time the security interest was created, the secured party neither had knowledge of nor consented to the act or omission. No personal property may be forfeited under this subsection (1)(g), to the extent of the interest of an owner, by reason of any act or omission which that owner establishes was committed or omitted without the owner’s knowledge or consent; and
   (h) All real property, including any right, title, and interest in the whole of any lot or tract of land, and any appurtenances or improvements which are being used with the knowledge of the owner for the manufacturing, compounding, processing, delivery, importing, or exporting of any controlled substance, or which have been acquired in whole or in part with proceeds traceable to an exchange or series of exchanges in violation of this chapter or chapter 69.41 or 69.52 RCW, if such activity is not less than a class C felony and a substantial nexus exists between the commercial production or sale of the controlled substance and the real property. However:
      (i) No property may be forfeited pursuant to this subsection (1)(h), to the extent of the interest of an owner, by reason
of any act or omission committed or omitted without the owner’s knowledge or consent;

(ii) The bona fide gift of a controlled substance, legend drug, or imitation controlled substance shall not result in the forfeiture of real property;

(iii) The possession of marijuana shall not result in the forfeiture of real property unless the marijuana is possessed for commercial purposes, the amount possessed is five or more plants or one pound or more of marijuana, and a substantial nexus exists between the possession of marijuana and the real property. In such a case, the intent of the offender shall be determined by the preponderance of the evidence, including the offender’s prior criminal history, the amount of marijuana possessed by the offender, the sophistication of the activity or equipment used by the offender, and other evidence which demonstrates the offender’s intent to engage in commercial activity;

(iv) The unlawful sale of marijuana or a legend drug shall not result in the forfeiture of real property unless the sale was forty grams or more in the case of marijuana or one hundred dollars or more in the case of a legend drug, and a substantial nexus exists between the unlawful sale and the real property; and

(v) A forfeiture of real property encumbered by a bona fide security interest is subject to the interest of the secured party if the secured party, at the time the security interest was created, neither had knowledge of nor consented to the act or omission.

(2) Real or personal property subject to forfeiture under this chapter may be seized by any board inspector or law enforcement officer of this state upon process issued by any superior court having jurisdiction over the property. Seizure of real property shall include the filing of a lis pendens by the seizing agency. Real property seized under this section shall not be transferred or otherwise conveyed until ninety days after seizure or until a judgment of forfeiture is entered, whichever is later: PROVIDED, That real property seized under this section may be transferred or conveyed to any person or entity who acquires title by foreclosure or deed in lieu of foreclosure of a security interest. Seizure of personal property without process may be made if:

(a) The seizure is incident to an arrest or a search under a search warrant or an inspection under an administrative inspection warrant;

(b) The property subject to seizure has been the subject of a prior judgment in favor of the state in a criminal injunction or forfeiture proceeding based upon this chapter;

(c) A board inspector or law enforcement officer has probable cause to believe that the property is directly or indirectly dangerous to health or safety; or

(d) The board inspector or law enforcement officer has probable cause to believe that the property was used or is intended to be used in violation of this chapter.

(3) In the event of seizure pursuant to subsection (2) of this section, proceedings for forfeiture shall be deemed commenced by the seizure. The law enforcement agency under whose authority the seizure was made shall cause notice to be served within fifteen days following the seizure on the owner of the property seized and the person in charge thereof and any person having any known right or interest therein, including any community property interest, of the seizure and intended forfeiture of the seized property. Service of notice of seizure of real property shall be made according to the rules of civil procedure. However, the state may not obtain a default judgment with respect to real property against a party who is served by substituted service absent an affidavit stating that a good faith effort has been made to ascertain if the defaulted party is incarcerated within the state, and that there is no present basis to believe that the party is incarcerated within the state. Notice of seizure in the case of property subject to a security interest that has been perfected by filing a financing statement in accordance with chapter 62A.9A RCW, or a certificate of title, shall be made by service upon the secured party or the secured party’s assignee at the address shown on the financing statement or the certificate of title. The notice of seizure in other cases may be served by any method authorized by law or court rule including but not limited to service by certified mail with return receipt requested. Service by mail shall be deemed complete upon mailing within the fifteen day period following the seizure.

(4) If no person notifies the seizing law enforcement agency in writing of the person’s claim of ownership or right to possession of items specified in subsection (1)(d), (g), or (h) of this section within forty-five days of the service of notice from the seizing agency in the case of personal property and ninety days in the case of real property, the item seized shall be deemed forfeited. The community property interest in real property of a person whose spouse or domestic partner committed a violation giving rise to seizure of the real property may not be forfeited if the person did not participate in the violation.

(5) If any person notifies the seizing law enforcement agency in writing of the person’s claim of ownership or right to possession of items specified in subsection (1)(b), (c), (d), (e), (f), (g), or (h) of this section within forty-five days of the service of notice from the seizing agency in the case of personal property and ninety days in the case of real property, the person or persons shall be afforded a reasonable opportunity to be heard as to the claim or right. The notice of claim may be served by any method authorized by law or court rule including, but not limited to, service by first-class mail. Service by mail shall be deemed complete upon mailing within the forty-five day period following service of the notice of seizure in the case of personal property and within the ninety-day period following service of the notice of seizure in the case of real property. The hearing shall be before the chief law enforcement officer of the seizing agency or the chief law enforcement officer’s designee, except where the seizing agency is a state agency as defined in RCW 34.12.020(4), the hearing shall be before the chief law enforcement officer of the seizing agency or an administrative law judge appointed under chapter 34.12 RCW, except that any person asserting a claim or right may remove the matter to a court of competent jurisdiction. Removal of any matter involving personal property may only be accomplished according to the rules of civil procedure. The person seeking removal of the matter must serve process against the state, county, political subdivision, or municipality that operates the seizing agency, and any other party of interest, in accordance with RCW 4.28.080 or 4.92.020, within forty-five days after the person seeking removal has notified the seizing law enforcement agency of the person’s claim of ownership or right to possession. The
court to which the matter is to be removed shall be the district court when the aggregate value of personal property is within the jurisdictional limit set forth in RCW 3.66.020. A hearing before the seizing agency and any appeal therefrom shall be under Title 34 RCW. In all cases, the burden of proof is upon the law enforcement agency to establish, by a preponderance of the evidence, that the property is subject to forfeiture.

The seizing law enforcement agency shall promptly return the article or articles to the claimant upon a determination by the administrative law judge or court that the claimant is the present lawful owner or is lawfully entitled to possession thereof of items specified in subsection (1)(b), (c), (d), (e), (f), (g), or (h) of this section.

(6) In any proceeding to forfeit property under this title, where the claimant substantially prevails, the claimant is entitled to reasonable attorneys’ fees reasonably incurred by the claimant. In addition, in a court hearing between two or more claimants to the article or articles involved, the prevailing party is entitled to a judgment for costs and reasonable attorneys’ fees.

(7) When property is forfeited under this chapter the board or seizing law enforcement agency may:

(a) Retain it for official use or upon application by any law enforcement agency of this state release such property to such agency for the exclusive use of enforcing the provisions of this chapter;

(b) Sell that which is not required to be destroyed by law and which is not harmful to the public;

(c) Request the appropriate sheriff or director of public safety to take custody of the property and remove it for disposition in accordance with law; or

(d) Forward it to the drug enforcement administration for disposition.

(8)(a) When property is forfeited, the seizing agency shall keep a record indicating the identity of the prior owner, if known, a description of the property, the disposition of the property, the value of the property at the time of seizure, and the amount of proceeds realized from disposition of the property.

(b) Each seizing agency shall retain records of forfeited property for at least seven years.

(c) Each seizing agency shall file a report including a copy of the records of forfeited property with the state treasurer each calendar quarter.

(d) The quarterly report need not include a record of forfeited property that is still being held for use as evidence during the investigation or prosecution of a case or during the appeal from a conviction.

(9)(a) By January 31st of each year, each seizing agency shall remit to the state treasurer an amount equal to ten percent of the net proceeds of any property forfeited during the preceding calendar year. Money remitted shall be deposited in the state general fund.

(b) The net proceeds of forfeited property is the value of the forfeitable interest in the property after deducting the cost of satisfying any bona fide security interest to which the property is subject at the time of seizure; and in the case of sold property, after deducting the cost of sale, including reasonable fees or commissions paid to independent selling agents, and the cost of any valid landlord’s claim for damages under subsection (15) of this section.

(c) The value of sold forfeited property is the sale price. The value of retained forfeited property is the fair market value of the property at the time of seizure, determined when possible by reference to an applicable commonly used index, such as the index used by the department of licensing for valuation of motor vehicles. A seizing agency may use, but need not use, an independent qualified appraiser to determine the value of retained property. If an appraiser is used, the value of the property appraised is net of the cost of the appraisal. The value of destroyed property and retained firearms or illegal property is zero.

(10) Forfeited property and net proceeds not required to be paid to the state treasurer shall be retained by the seizing law enforcement agency exclusively for the expansion and improvement of controlled substances related law enforcement activity. Money retained under this section may not be used to supplant preexisting funding sources.

(11) Controlled substances listed in Schedule I, II, III, IV, and V that are possessed, transferred, sold, or offered for sale in violation of this chapter are contraband and shall be seized and summarily forfeited to the state. Controlled substances listed in Schedule I, II, III, IV, and V, which are seized or come into the possession of the board, the owners of which are unknown, are contraband and shall be summarily forfeited to the board.

(12) Species of plants from which controlled substances in Schedules I and II may be derived which have been planted or cultivated in violation of this chapter, or of which the owners or cultivators are unknown, or which are wild growths, may be seized and summarily forfeited to the board.

(13) The failure, upon demand by a board inspector or law enforcement officer, of the person in occupancy or in control of land or premises upon which the species of plants are growing or being stored to produce an appropriate registration or proof that he or she is the holder thereof constitutes authority for the seizure and forfeiture of the plants.

(14) Upon the entry of an order of forfeiture of real property, the court shall forward a copy of the order to the assessor of the county in which the property is located. Orders for the forfeiture of real property shall be entered by the superior court, subject to court rules. Such an order shall be filed by the seizing agency in the county auditor’s records in the county in which the real property is located.

(15) A landlord may assert a claim against proceeds from the sale of assets seized and forfeited under subsection (7)(b) of this section, only if:

(a) A law enforcement officer, while acting in his or her official capacity, directly caused damage to the complaining landlord’s property while executing a search of a tenant’s residence; and

(b) The landlord has applied any funds remaining in the tenant’s deposit, to which the landlord has a right under chapter 59.18 RCW, to cover the damage directly caused by a law enforcement officer prior to asserting a claim under the provisions of this section;

(i) Only if the funds applied under (b) of this subsection are insufficient to satisfy the damage directly caused by a law enforcement officer, may the landlord seek compensation for the damage by filing a claim against the governmental entity under whose authority the law enforcement agency operates within thirty days after the search;
(ii) Only if the governmental entity denies or fails to respond to the landlord’s claim within sixty days of the date of filing, may the landlord collect damages under this subsection by filing within thirty days of denial or the expiration of the sixty-day period, whichever occurs first, a claim with the seizing law enforcement agency. The seizing law enforcement agency must notify the landlord of the status of the claim by the end of the thirty-day period. Nothing in this section requires the claim to be paid by the end of the sixty-day or thirty-day period.

(c) For any claim filed under (b) of this subsection, the law enforcement agency shall pay the claim unless the agency provides substantial proof that the landlord either:

(i) Knew or consented to actions of the tenant in violation of this chapter or chapter 69.41 or 69.52 RCW; or

(ii) Failed to respond to a notification of the illegal activity, provided by a law enforcement agency under RCW 59.18.075, within seven days of receipt of notification of the illegal activity.

(16) The landlord’s claim for damages under subsection (15) of this section may not include a claim for loss of business and is limited to:

(a) Damage to tangible property and clean-up costs;

(b) The lesser of the cost of repair or fair market value of the damage directly caused by a law enforcement officer;

(c) The proceeds from the sale of the specific tenant’s property seized and forfeited under subsection (7)(b) of this section; and

(d) The proceeds available after the seizing law enforcement agency satisfies any bona fide security interest in the tenant’s property and costs related to sale of the tenant’s property as provided by subsection (9)(b) of this section.

(17) Subsections (15) and (16) of this section do not limit any other rights a landlord may have against a tenant to collect for damages. However, if a law enforcement agency satisfies a landlord’s claim under subsection (15) of this section, the rights the landlord has against the tenant for damages directly caused by a law enforcement officer under the terms of the landlord and tenant’s contract are subrogated to the law enforcement agency. [2009 c 479 § 46; 2009 c 364 § 1; 2008 c 6 § 631; 2003 c 53 § 348; 2001 c 168 § 1; 1993 c 487 § 1; 1992 c 211 § 1. Prior: 1992 c 210 § 5 repealed by 1992 c 211 § 2); 1990 c 248 § 2; 1990 c 213 § 12; 1989 c 271 § 212; 1988 c 282 § 2; 1986 c 124 § 9; 1984 c 258 § 333; 1983 c 2 § 15; prior: 1982 c 189 § 6; 1982 c 171 § 1; prior: 1981 c 67 § 32; 1981 c 48 § 3; 1977 ex.s. c 77 § 1; 1971 ex.s. c 308 § 69.50.505.]

Reviser’s note: This section was amended by 2009 c 364 § 1 and by 2009 c 479 § 46, each without reference to the other. Both amendments are incorporated in the publication of this section under RCW 1.12.025(2). For rule of construction, see RCW 1.12.025(1).

Effective date—2009 c 479: See note following RCW 2.56.030.

Part headings not law—Severability—2008 c 6: See RCW 26.60.900 and 26.60.901.

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

Findings—1989 c 271: "The legislature finds that: Drug offenses and crimes resulting from illegal drug use are destructive to society; the nature of drug trafficking results in many property crimes and crimes of violence; state and local governmental agencies incur immense expenses in the investigation, prosecution, adjudication, incarceration, and treatment of drug-related offenders and the compensation of their victims; drug-related offenses are difficult to eradicate because of the profits derived from the criminal activity, which can be invested in legitimate assets and later used for further criminal activities; and the forfeiture of real assets where a substantial nexus exists between the commercial production or sale of the substances and the real property will provide a significant deterrent to crime by removing the profit incentive of drug trafficking, and will provide a revenue source that will partially defray the large costs incurred by government as a result of these crimes. The legislature recognizes that seizure of real property is a very powerful tool and should not be applied in cases in which a manifest injustice would occur as a result of forfeiture of an innocent spouse’s community property interest." [1989 c 271 § 211.]

Intent—1984 c 258: See note following RCW 3.34.130.

Additional notes found at www.leg.wa.gov

69.50.506 Burden of proof; liabilities. (a) It is not necessary for the state to negate any exemption or exception in this chapter in any complaint, information, indictment, or other pleading or in any trial, hearing, or other proceeding under this chapter. The burden of proof of any exemption or exception is upon the person claiming it.

(b) In the absence of proof that a person is the duly authorized holder of an appropriate registration or order form issued under this chapter, he or she is presumed not to be the holder of the registration or form. The burden of proof is upon him or her to rebut the presumption.

(c) No liability is imposed by this chapter upon any authorized state, county, or municipal officer, engaged in the lawful performance of his or her duties. [2012 c 117 § 371; 1971 ex.s. c 308 § 69.50.506.]

69.50.507 Judicial review. All final determinations, findings, and conclusions of the state board of pharmacy under this chapter are final and conclusive decisions of the matters involved. Any person aggrieved by the decision may obtain review of the decision in the superior court wherein he or she resides or in the superior court of Thurston county, such review to be in conformity with the administrative procedure act, chapter 34.05 RCW. [2012 c 117 § 371; 1971 ex.s. c 308 § 69.50.507.]

69.50.508 Education and research. (a) The state board of pharmacy may carry out educational programs designed to prevent and deter misuse and abuse of controlled substances. In connection with these programs it may:

(1) promote better recognition of the problems of misuse and abuse of controlled substances within the regulated industry and among interested groups and organizations;

(2) assist the regulated industry and interested groups and organizations in contributing to the reduction of misuse and abuse of controlled substances;

(3) consult with interested groups and organizations to aid them in solving administrative and organizational problems;

(4) evaluate procedures, projects, techniques, and controls conducted or proposed as part of educational programs on misuse and abuse of controlled substances;

(5) disseminate the results of research on misuse and abuse of controlled substances to promote a better public understanding of what problems exist and what can be done to combat them; and

(6) assist in the education and training of state and local law enforcement officials in their efforts to control misuse and abuse of controlled substances.
(b) The board may encourage research on misuse and abuse of controlled substances. In connection with the research, and in furtherance of the enforcement of this chapter, it may:

1. establish methods to assess accurately the effects of controlled substances and identify and characterize those with potential for abuse;
2. make studies and undertake programs of research to:
   i. develop new or improved approaches, techniques, systems, equipment and devices to strengthen the enforcement of this chapter;
   ii. determine patterns of misuse and abuse of controlled substances and the social effects thereof; and,
   iii. improve methods for preventing, predicting, understanding and dealing with the misuse and abuse of controlled substances; and,
3. enter into contracts with public agencies, institutions of higher education, and private organizations or individuals for the purpose of conducting research, demonstrations, or special projects which bear directly on misuse and abuse of controlled substances.

(c) The board may enter into contracts for educational and research activities without performance bonds.

(d) The board may authorize persons engaged in research on the use and effects of controlled substances to withhold the names and other identifying characteristics of individuals who are the subjects of the research. Persons who obtain this authorization are not compelled in any civil, criminal, administrative, legislative, or other proceeding to identify the individuals who are the subjects of research for which the authorization was obtained.

(e) The board may authorize the possession and distribution of controlled substances by persons engaged in research. Persons who obtain this authorization are exempt from state prosecution for possession and distribution of controlled substances to the extent of the authorization. [1971 ex.s. c 308 § 69.50.508.]

69.50.509 Search and seizure of controlled substances. If, upon the sworn complaint of any person, it shall be made to appear to any judge of the superior court, district court, or municipal court that there is probable cause to believe that any controlled substance is being used, manufactured, sold, bartered, exchanged, administered, dispensed, delivered, distributed, produced, possessed, given away, furnished or otherwise disposed of or kept in violation of the provisions of this chapter, such judge shall, with or without the approval of the prosecuting attorney, issue a warrant directed to any law enforcement officer of the state, commanding him or her to search the premises designated and described in such complaint and warrant, and to seize all controlled substances there found, together with the vessels in which they are contained, and all implements, furniture and fixtures used or kept for the illegal manufacture, sale, barter, exchange, administering, dispensing, delivering, distributing, producing, possessing, giving away, furnishing or otherwise disposing of such controlled substances, and to safely keep the same, and to make a return of said warrant within three days, showing all acts and things done thereunder, with a particular statement of all articles seized and the name of the person or persons in whose possession the same were found, if any, and if no person be found in the possession of said articles, the returns shall so state. The provisions of RCW 10.31.030 as now or hereafter amended shall apply to actions taken pursuant to this chapter. [1987 c 202 § 228; 1971 ex.s. c 308 § 69.50.509.]

Intent—1987 c 202: See note following RCW 2.04.190.

69.50.510 Search and seizure at rental premises—Notification of landlord. Whenever a controlled substance which is manufactured, distributed, dispensed, or acquired in violation of this chapter is seized at rental premises, the law enforcement agency shall make a reasonable attempt to discover the identity of the landlord and shall notify the landlord in writing, at the last address listed in the property tax records and at any other address known by the law enforcement agency, of the seizure and the location of the seizure. [1988 c 150 § 9.]

Legislative findings—Severability—1988 c 150: See notes following RCW 59.18.130.

69.50.511 Cleanup of hazardous substances at illegal drug manufacturing facility—Rules. Law enforcement agencies who during the official investigation or enforcement of any illegal drug manufacturing facility come in contact with or are aware of any substances suspected of being hazardous as defined in RCW 70.105D.020, shall notify the department of ecology for the purpose of securing a contractor to identify, clean up, store, and dispose of suspected hazardous substances, except for those random and representative samples obtained for evidentiary purposes. Whenever possible, a destruct order covering hazardous substances which may be described in general terms shall be obtained concurrently with a search warrant. Materials that have been photographed, fingerprinted, and subsampled by police shall be destroyed as soon as practical. The department of ecology shall make every effort to recover costs from the parties responsible for the suspected hazardous substance. All recoveries shall be deposited in the account or fund from which contractor payments are made.

The department of ecology may adopt rules to carry out its responsibilities under this section. The department of ecology shall consult with law enforcement agencies prior to adopting any rule or policy relating to this section. [2007 c 104 § 17; 1990 c 213 § 13; 1989 c 271 § 228.]

Application—Construction—Severability—2007 c 104: See RCW 64.70.015 and 64.70.900.

Additional notes found at www.leg.wa.gov

69.50.525 Diversion prevention and control—Report. (a) As used in this section, "diversion" means the transfer of any controlled substance from a licit to an illicit channel of distribution or use.
(b) The department shall regularly prepare and make available to other state regulatory, licensing, and law enforcement agencies a report on the patterns and trends of actual distribution, diversion, and abuse of controlled substances.
(c) The department shall enter into written agreements with local, state, and federal agencies for the purpose of improving identification of sources of diversion and to improve enforcement of and compliance with this chapter and other laws and regulations pertaining to unlawful con-
duct involving controlled substances. An agreement must specify the roles and responsibilities of each agency that has information or authority to identify, prevent, and control drug diversions and drug abuse. The department shall convene periodic meetings to coordinate a state diversion prevention and control program. The department shall arrange for cooperation and exchange of information among agencies and with neighboring states and the federal government. [1998 c 245 § 109; 1993 c 187 § 20.]

ARTICLE VI
MISCELLANEOUS

69.50.601 Pending proceedings. (a) Prosecution for any violation of law occurring prior to May 21, 1971 is not affected or abated by this chapter. If the offense being prosecuted is similar to one set out in Article IV of this chapter, then the penalties under Article IV apply if they are less than those under prior law.

(b) Civil seizures or forfeitures and injunctive proceedings commenced prior to May 21, 1971 are not affected by this chapter.

(c) All administrative proceedings pending under prior laws which are superseded by this chapter shall be continued and brought to a final determination in accord with the laws and rules in effect prior to May 21, 1971. Any substance controlled under prior law which is not listed within Schedules I through V, is automatically controlled without further proceedings and shall be listed in the appropriate schedule.

(d) The state board of pharmacy shall initially permit persons to register who own or operate any establishment engaged in the manufacture, distribution, or dispensing of any controlled substance prior to May 21, 1971 and who are registered or licensed by the state.

(e) This chapter applies to violations of law, seizures and forfeiture, injunctive proceedings, administrative proceedings and investigations which occur following May 21, 1971. [1971 ex.s. c 308 § 69.50.601.]

69.50.602 Continuation of rules. Any orders and rules promulgated under any law affected by this chapter and in effect on May 21, 1971 and not in conflict with it continue in effect until modified, superseded or repealed. [1971 ex.s. c 308 § 69.50.602.]

69.50.603 Uniformity of interpretation. This chapter shall be so applied and construed as to effectuate its general purpose to make uniform the law with respect to the subject of this chapter among those states which enact it. [1971 ex.s. c 308 § 69.50.603.]

69.50.604 Short title. This chapter may be cited as the Uniform Controlled Substances Act. [1971 ex.s. c 308 § 69.50.604.]

69.50.605 Severability—1971 ex.s. c 308. If any provision of this act or the application thereof to any person or circumstance is held invalid, the invalidity does not affect other provisions or applications of the act which can be given effect without the invalid provision or application, and to this end the provisions of this act are severable. [1971 ex.s. c 308 § 69.50.605.]

69.50.606 Repealers. The laws specified below are repealed except with respect to rights and duties which matured, penalties which were incurred and proceedings which were begun before the effective date of this act:

(1) Section 2072, Code of 1881, section 418, chapter 249, Laws of 1909, section 4, chapter 205, Laws of 1963 and RCW 9.91.030;

(2) Section 69.33.220, chapter 27, Laws of 1959, section 7, chapter 256, Laws of 1969 ex. sess. and RCW 69.33.220;

(3) Sections 69.33.230 through 69.33.280, chapter 27, Laws of 1959 and RCW 69.33.230 through 69.33.280;

(4) Section 69.33.290, chapter 27, Laws of 1959, section 1, chapter 97, Laws of 1959 and RCW 69.33.290;

(5) Section 69.33.300, chapter 27, Laws of 1959, section 8, chapter 256, Laws of 1969 ex. sess. and RCW 69.33.300;

(6) Sections 69.33.310 through 69.33.400, chapter 27, Laws of 1959 and RCW 69.33.310 through 69.33.400;

(7) Section 69.33.410, chapter 27, Laws of 1959, section 20, chapter 38, Laws of 1963 and RCW 69.33.410;

(8) Sections 69.33.420 through 69.33.440, 69.33.900 through 69.33.950, chapter 27, Laws of 1959 and RCW 69.33.420 through 69.33.440, 69.33.900 through 69.33.950;

(9) Section 255, chapter 249, Laws of 1909 and RCW 69.40.040;

(10) Section 1, chapter 6, Laws of 1939, section 1, chapter 29, Laws of 1939, section 1, chapter 57, Laws of 1945, section 1, chapter 24, Laws of 1955, section 1, chapter 49, Laws of 1961, section 1, chapter 71, Laws of 1967, section 9, chapter 256, Laws of 1969 ex. sess. and RCW 69.40.060;


(12) Section 21, chapter 38, Laws of 1963 and RCW 69.40.063;


(14) Section 12, chapter 256, Laws of 1969 ex. sess. and RCW 69.40.075;

(15) Section 1, chapter 205, Laws of 1963 and RCW 69.40.080;

(16) Section 2, chapter 205, Laws of 1963 and RCW 69.40.090;

(17) Section 3, chapter 205, Laws of 1963 and RCW 69.40.100;

(18) Section 11, chapter 256, Laws of 1969 ex. sess. and RCW 69.40.110;

(19) Section 1, chapter 33, Laws of 1970 ex. sess. and RCW 69.40.120; and

(20) Section 1, chapter 80, Laws of 1970 ex. sess. [1971 ex.s. c 308 § 69.50.606.]

69.50.607 Effective date—1971 ex.s. c 308. This act is necessary for the immediate preservation of the public peace, health and safety, the support of the state government and its existing public institutions, and shall take effect immediately. [1971 ex.s. c 308 § 69.50.607.]
69.50.608 State preemption. The state of Washington fully occupies and preempts the entire field of setting penalties for violations of the controlled substances act. Cities, towns, and counties or other municipalities may enact only those laws and ordinances relating to controlled substances that are consistent with this chapter. Such local ordinances shall have the same penalties as provided for by state law. Local laws and ordinances that are inconsistent with the requirements of state law shall not be enacted and are preempted and repealed, regardless of the nature of the code, charter, or home rule status of the city, town, county, or municipality. [1989 c 271 § 601.]

69.50.609 Captions not law—1993 c 187. Section captions as used in this act constitute no part of the law. [1993 c 187 § 23.]

Chapter 69.51 RCW
CONTROLLED SUBSTANCES THERAPEUTIC RESEARCH ACT

Sections
69.51.010 Short title. This chapter may be cited as the Controlled Substances Therapeutic Research Act. [1979 c 136 § 1.]

69.51.020 Legislative purpose. The legislature finds that recent research has shown that the use of marijuana may alleviate the nausea and ill effects of cancer chemotherapy and radiotherapy, and, additionally, may alleviate the ill effects of glaucoma. The legislature further finds that there is a need for further research and experimentation regarding the use of marijuana under strictly controlled circumstances. It is for this purpose that the Controlled Substances Therapeutic Research Act is hereby enacted. [1979 c 136 § 2.]

69.51.030 Definitions. As used in this chapter:
(1) "Board" means the state board of pharmacy;
(2) "Department" means the department of health;
(3) "Marijuana" means all parts of the plant of the genus Cannabis L., whether growing or not, the seeds thereof, the resin extracted from any part of the plant, and every compound, manufacture, salt, derivative, mixture, or preparation of the plant, its seeds, or resin; and
(4) "Practitioner" means a physician licensed pursuant to chapter 18.71 or 18.57 RCW. [1989 1st ex.s. c 9 § 438; 1979 c 136 § 3.]

69.51.040 Controlled substances therapeutic research program. (1) There is established in the board the controlled substances therapeutic research program. The program shall be administered by the department. The board shall promulgate rules necessary for the proper administration of the Controlled Substances Therapeutic Research Act. In such promulgation, the board shall take into consideration those pertinent rules promulgated by the United States drug enforcement agency, the food and drug administration, and the national institute on drug abuse.

(2) Except as provided in RCW 69.51.050(4), the controlled substances therapeutic research program shall be limited to cancer chemotherapy and radiology patients and glaucoma patients, who are certified to the patient qualification review committee as being involved in a life-threatening or sense-threatening situation. No patient may be admitted to the controlled substances therapeutic research program without full disclosure by the practitioner of the experimental nature of this program and of the possible risks and side effects of the proposed treatment in accordance with the informed consent provisions of chapter 7.70 RCW.

(3) The board shall provide by rule for a program of registration with the department of bona fide controlled substance therapeutic research projects. [1989 1st ex.s. c 9 § 439; 1979 c 136 § 4.]

Additional notes found at www.leg.wa.gov

69.51.050 Patient qualification review committee. (1) The board shall appoint a patient qualification review committee to serve at its pleasure. The patient qualification review committee shall be comprised of:
(a) A physician licensed to practice medicine in Washington state and specializing in the practice of ophthalmology;
(b) A physician licensed to practice medicine in Washington state and specializing in the subspecialty of medical oncology;
(c) A physician licensed to practice medicine in Washington state and specializing in the practice of psychiatry; and
(d) A physician licensed to practice medicine in Washington state and specializing in the practice of radiology.

Members of the committee shall be compensated at the rate of fifty dollars per day for each day spent in the performance of their official duties, and shall receive reimbursement for their travel expenses as provided in RCW 43.03.050 and 43.03.060.

(2) The patient qualification review committee shall review all applicants for the controlled substance therapeutic research program and their licensed practitioners and certify their participation in the program.

(3) The patient qualification review committee and the board shall insure that the privacy of individuals who participate in the controlled substance therapeutic research program is protected by withholding from all persons not connected with the conduct of the research the names and other identifying characteristics of such individuals. Persons authorized to engage in research under the controlled substance therapeutic research program may not be compelled in any civil, criminal, administrative, legislative, or other proceeding to identify the individuals who are the subjects of research for which the authorization was granted, except to the extent necessary to permit the board to determine whether the research is being conducted in accordance with the authorization.

(4) The patient qualification review committee may include other disease groups for participation in the controlled substances therapeutic research program after perti-
69.51.060 Sources and distribution of marijuana. (1) The board shall obtain marijuana through whatever means it deems most appropriate and consistent with regulations promulgated by the United States food and drug administration, the drug enforcement agency, and the national institute on drug abuse. [1979 c 136 § 5.]

(2) The board may use marijuana which has been confiscated by local or state law enforcement agencies and has been determined to be free from contamination.

(3) The board shall distribute the analyzed marijuana to approved practitioners and/or institutions in accordance with rules promulgated by the board. [1979 c 136 § 6.]

69.51.080 Cannabis and related products considered Schedule II substances. (1) The enumeration of tetrahydrocannabinols, or a chemical derivative of tetrahydrocannabinols in RCW 69.50.204 as a Schedule I controlled substance does not apply to the use of cannabis, tetrahydrocannabinols, or a chemical derivative of tetrahydrocannabinols by certified patients pursuant to the provisions of this chapter.

(2) Cannabis, tetrahydrocannabinols, or a chemical derivative of tetrahydrocannabinols shall be considered Schedule II substances as enumerated in RCW 69.50.206 only for the purposes enumerated in this chapter. [1979 c 136 § 8.]

## Chapter 69.51A RCW

### MEDICAL CANNABIS

(Formerly: Medical marijuana)

Sections

- 69.51A.005 Purpose and intent.
- 69.51A.010 Definitions.
- 69.51A.020 Construction of chapter.
- 69.51A.025 Construction of chapter—Compliance with RCW 69.51A.040.
- 69.51A.030 Acts not constituting crimes or unprofessional conduct—Health care professionals not subject to penalties or liabilities.
- 69.51A.040 Compliance with chapter—Qualifying patients and designated providers not subject to penalties—Law enforcement not subject to liability.
- 69.51A.043 Failure to register—Affirmative defense.
- 69.51A.045 Possession of cannabis exceeding lawful amount—Affirmative defense.
- 69.51A.047 Failure to register or present valid documentation—Affirmative defense.
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69.51A.902 Captions not law—1999 c 2.
69.51A.903 Severability—2011 c 181.

### 69.51A.005 Purpose and intent.

(1) The legislature finds that:

(a) There is medical evidence that some patients with terminal or debilitating medical conditions may, under their health care professional’s care, benefit from the medical use of cannabis. Some of the conditions for which cannabis appears to be beneficial include, but are not limited to:

(i) Nausea, vomiting, and cachexia associated with cancer, HIV-positive status, AIDS, hepatitis C, anorexia, and their treatments;

(ii) Severe muscle spasms associated with multiple sclerosis, epilepsy, and other seizure and spasticity disorders;

(iii) Acute or chronic glaucoma;

(iv) Crohn’s disease; and

(v) Some forms of intractable pain.

(b) Humanitarian compassion necessitates that the decision to use cannabis by patients with terminal or debilitating medical conditions is a personal, individual decision, based upon their health care professional’s professional medical judgment and discretion.

(2) Therefore, the legislature intends that:

(a) Qualifying patients with terminal or debilitating medical conditions who, in the judgment of their health care professionals, may benefit from the medical use of cannabis, shall not be arrested, prosecuted, or subject to other criminal sanctions or civil consequences under state law based solely on their medical use of cannabis, notwithstanding any other provision of law;

(b) Persons who act as designated providers to such patients shall also not be arrested, prosecuted, or subject to other criminal sanctions or civil consequences under state law, notwithstanding any other provision of law, based solely on their assisting with the medical use of cannabis; and

(c) Health care professionals shall also not be arrested, prosecuted, or subject to other criminal sanctions or civil consequences under state law for the proper authorization of medical use of cannabis by qualifying patients for whom, in the health care professional’s professional judgment, the medical use of cannabis may prove beneficial.

(3) Nothing in this chapter establishes the medical necessity or medical appropriateness of cannabis for treating terminal or debilitating medical conditions as defined in RCW 69.51A.010.

(4) Nothing in this chapter diminishes the authority of correctional agencies and departments, including local governments or jails, to establish a procedure for determining when the use of cannabis would impact community safety or the effective supervision of those on active supervision for a criminal conviction, nor does it create the right to any accommodation of any medical use of cannabis in any correctional facility or jail. [2011 c 181 § 102; 2010 c 284 § 1; 2007 c 371 § 2; 1999 c 2 § 2 (Initiative Measure No. 692, approved November 3, 1998).]

### Intent—2007 c 371:

"The legislature intends to clarify the law on medical marijuana so that the lawful use of this substance is not impaired and medical practitioners are able to exercise their best professional judgment in the delivery of medical treatment, qualifying patients may fully participate in the medical use of marijuana, and designated providers may assist patients in the manner provided by this act without fear of state criminal prosecution."
69.51A.010 Definitions. The definitions in this section apply throughout this chapter unless the context clearly requires otherwise.

(1) "Designated provider" means a person who:
   (a) Is eighteen years of age or older;
   (b) Has been designated in writing by a patient to serve as a designated provider under this chapter;
   (c) Is prohibited from consuming marijuana obtained for personal, medical use of the patient for whom the individual is acting as designated provider; and
   (d) Is the designated provider to only one patient at any one time.

(2) "Health care professional," for purposes of this chapter, means a physician licensed under chapter 18.71 RCW, a physician assistant licensed under chapter 18.71A RCW, an osteopathic physician licensed under chapter 18.57 RCW, an osteopathic physicians’ assistant licensed under chapter 18.57A RCW, a naturopath licensed under chapter 18.36A RCW, or an advanced registered nurse practitioner licensed under chapter 18.79 RCW.

(3) "Medical use of marijuana" means the production, possession, or administration of marijuana, as defined in *RCW 69.50.101(a), for the exclusive benefit of a qualifying patient in the treatment of his or her terminal or debilitating illness.

(4) "Qualifying patient" means a person who:
   (a) Is a patient of a health care professional;
   (b) Has been diagnosed by that health care professional as having a terminal or debilitating medical condition;
   (c) Is a resident of the state of Washington at the time of such diagnosis;
   (d) Has been advised by that health care professional about the risks and benefits of the medical use of marijuana; and
   (e) Has been advised by that health care professional that they may benefit from the medical use of marijuana.

(5) "Tamper-resistant paper" means paper that meets one or more of the following industry-recognized features:
   (a) One or more features designed to prevent copying of the paper;
   (b) One or more features designed to prevent the erasure or modification of information on the paper; or
   (c) One or more features designed to prevent the use of counterfeit valid documentation.

(6) "Terminal or debilitating medical condition" means:
   (a) Cancer, human immunodeficiency virus (HIV), multiple sclerosis, epilepsy or other seizure disorder, or spasticity disorders; or
   (b) Intractable pain, limited for the purpose of this chapter to mean pain unrelieved by standard medical treatments and medications; or
   (c) Glaucome, either acute or chronic, limited for the purpose of this chapter to mean increased intraocular pressure unrelieved by standard treatments and medications; or
   (d) Crohn’s disease with debilitating symptoms unrelieved by standard treatments or medications; or
   (e) Hepatitis C with debilitating nausea or intractable pain unrelieved by standard treatments or medications; or
   (f) Diseases, including anorexia, which result in nausea, vomiting, wasting, appetite loss, cramping, seizures, muscle spasms, or spasticity, when these symptoms are unrelieved by standard treatments or medications; or
   (g) Any other medical condition duly approved by the Washington state medical quality assurance commission in consultation with the board of osteopathic medicine and surgery as directed in this chapter.

(7) "Valid documentation" means:
   (a) A statement signed and dated by a qualifying patient’s health care professional written on tamper-resistant paper, which states that, in the health care professional’s professional opinion, the patient may benefit from the medical use of marijuana; and
   (b) Proof of identity such as a Washington state driver’s license or identicard, as defined in RCW 46.20.035. [2010 c 284 § 2; 2007 c 371 § 3; 1999 c 2 § 6 (Initiative Measure No. 692, approved November 3, 1998).]

*Reviser’s note: RCW 69.50.101 was alphabetized pursuant to RCW 1.08.015(2)(k), changing subsection (q) to subsection (r).

Intent—2007 c 371: See note following RCW 69.51A.005.

69.51A.020 Construction of chapter. Nothing in this chapter shall be construed to supersede Washington state law prohibiting the acquisition, possession, manufacture, sale, or use of cannabis for nonmedical purposes. Criminal penalties created under chapter 181, Laws of 2011 do not preclude the prosecution or punishment for other crimes, including other crimes involving the manufacture or delivery of cannabis for nonmedical purposes. [2011 c 181 § 103; 1999 c 2 § 3 (Initiative Measure No. 692, approved November 3, 1998).]

69.51A.025 Construction of chapter—Compliance with RCW 69.51A.040. Nothing in this chapter or in the rules adopted to implement it precludes a qualifying patient or designated provider from engaging in the private, unlicensed, noncommercial production, possession, transportation, delivery, or administration of cannabis for medical use as authorized under RCW 69.51A.040. [2011 c 181 § 413.]

69.51A.030 Acts not constituting crimes or unprofessional conduct—Health care professionals not subject to penalties or liabilities. (1) The following acts do not constitute crimes under state law or unprofessional conduct under chapter 18.130 RCW, and a health care professional may not be arrested, searched, prosecuted, disciplined, or subject to other criminal sanctions or civil consequences or liability under state law, or have real or personal property searched, seized, or forfeited pursuant to state law, notwithstanding any other provision of law as long as the health care professional complies with subsection (2) of this section:
   (a) Advising a patient about the risks and benefits of the medical use of cannabis or that the patient may benefit from the medical use of cannabis; or
   (b) Providing a patient meeting the criteria established under *RCW 69.51A.010(26) with valid documentation, based upon the health care professional’s assessment of the patient’s medical history and current medical condition, where such use is within a professional standard of care or in the individual health care professional’s medical judgment.
(2)(a) A health care professional may only provide a patient with valid documentation authorizing the medical use of cannabis or register the patient with the registry established in **section 901 of this act if he or she has a newly initiated or existing documented relationship with the patient, as a primary care provider or a specialist, relating to the diagnosis and ongoing treatment or monitoring of the patient’s terminal or debilitating medical condition, and only after:

(i) Completing a physical examination of the patient as appropriate, based on the patient’s condition and age;

(ii) Documenting the terminal or debilitating medical condition of the patient in the patient’s medical record and that the patient may benefit from treatment of this condition or its symptoms with medical use of cannabis;

(iii) Informing the patient of other options for treating the terminal or debilitating medical condition; and

(iv) Documenting other measures attempted to treat the terminal or debilitating medical condition that do not involve the medical use of cannabis.

(b) A health care professional shall not:

(i) Accept, solicit, or offer any form of pecuniary remuneration from or to a licensed dispenser, licensed producer, or licensed processor of cannabis products;

(ii) Offer a discount or any other thing of value to a qualifying patient who is a customer of, or agrees to be a customer of, a particular licensed dispenser, licensed producer, or licensed processor of cannabis products;

(iii) Examine or offer to examine a patient for purposes of diagnosing a terminal or debilitating medical condition at a location where cannabis is produced, processed, or dispensed;

(iv) Have a business or practice which consists solely of authorizing the medical use of cannabis;

(v) Include any statement or reference, visual or otherwise, on the medical use of cannabis in any advertisement for his or her business or practice; or

(vi) Hold an economic interest in an enterprise that produces, processes, or dispenses cannabis if the health care professional authorizes the medical use of cannabis.

(3) A violation of any provision of subsection (2) of this section constitutes unprofessional conduct under chapter 18.130 RCW. [2011 c 181 § 301; 2010 c 284 § 3; 2007 c 371 § 4; 1999 c 2 § 4 (Initiative Measure No. 692, approved November 3, 1998).]

Reviser’s note: *(1) RCW 69.51A.010(26) is a reference to the definition of "qualifying patient" which was amended and renumbered by 2011 c 181 § 201, but the section was vetoed by the governor.*

***(2) The section creating a registry, 2011 c 181 § 901, was vetoed by the governor.***

Intent—2007 c 371: See note following RCW 69.51A.005.

### 69.51A.040 Compliance with chapter—Qualifying patients and designated providers not subject to penalties—Law enforcement not subject to liability

The medical use of cannabis in accordance with the terms and conditions of this chapter does not constitute a crime and a qualifying patient or designated provider in compliance with the terms and conditions of this chapter may not be arrested, prosecuted, or subject to other criminal sanctions or civil consequences, for possession, manufacture, or delivery of, or for possession with intent to manufacture or deliver, cannabis under state law, or have real or personal property seized or forfeited for possession, manufacture, or delivery of, or for possession with intent to manufacture or deliver, cannabis under state law, and investigating peace officers and law enforcement agencies may not be held civilly liable for failure to seize cannabis in this circumstance, if:

(1)(a) The qualifying patient or designated provider possesses no more than fifteen cannabis plants and:

(i) No more than twenty-four ounces of usable cannabis;

(ii) No more cannabis product than what could reasonably be produced with no more than twenty-four ounces of usable cannabis; or

(iii) A combination of usable cannabis and cannabis product that does not exceed a combined total representing possession and processing of no more than twenty-four ounces of usable cannabis.

(b) If a person is both a qualifying patient and a designated provider for another qualifying patient, the person may possess no more than twice the amounts described in (a) of this subsection, whether the plants, usable cannabis, and cannabis product are possessed individually or in combination between the qualifying patient and his or her designated provider;

(2) The qualifying patient or designated provider presents his or her proof of registration with the department of health, to any peace officer who questions the patient or provider regarding his or her medical use of cannabis;

(3) The qualifying patient or designated provider keeps a copy of his or her proof of registration with the registry established in *section 901 of this act and the qualifying patient or designated provider’s contact information posted prominently next to any cannabis plants, cannabis products, or usable cannabis located at his or her residence;

(4) The investigating peace officer does not possess evidence that:

(a) The designated provider has converted cannabis produced or obtained for the qualifying patient for his or her own personal use or benefit; or

(b) The qualifying patient has converted cannabis produced or obtained for his or her own medical use to the qualifying patient’s personal, nonmedical use or benefit;

(5) The investigating peace officer does not possess evidence that the designated provider has served as a designated provider to more than one qualifying patient within a fifteen-day period; and

(6) The investigating peace officer has not observed evidence of any of the circumstances identified in *section 901(4) of this act. [2011 c 181 § 401; 2007 c 371 § 5; 1999 c 2 § 5 (Initiative Measure No. 692, approved November 3, 1998).]

*Reviser’s note: Section 901 of this act was vetoed by the governor.*

Intent—2007 c 371: See note following RCW 69.51A.005.

### 69.51A.043 Failure to register—Affirmative defense

(1) A qualifying patient or designated provider who is not registered with the registry established in *section 901 of this act may raise the affirmative defense set forth in subsection (2) of this section, if:

(a) The qualifying patient or designated provider presents his or her valid documentation to any peace officer who
questions the patient or provider regarding his or her medical use of cannabis;

(b) The qualifying patient or designated provider possesses no more cannabis than the limits set forth in RCW 69.51A.040(1);

(c) The qualifying patient or designated provider is in compliance with all other terms and conditions of this chapter;

(d) The investigating peace officer does not have probable cause to believe that the qualifying patient or designated provider has committed a felony, or is committing a misdemeanor in the officer’s presence, that does not relate to the medical use of cannabis;

(e) No outstanding warrant for arrest exists for the qualifying patient or designated provider; and

(f) The investigating peace officer has not observed evidence of any of the circumstances identified in *section 901(4) of this act.

(2) A qualifying patient or designated provider who is not registered with the registry established in *section 901 of this act, but who presents his or her valid documentation to any peace officer who questions the patient or provider regarding his or her medical use of cannabis, may assert an affirmative defense to charges of violations of state law relating to cannabis through proof at trial, by a preponderance of the evidence, that he or she otherwise meets the requirements of RCW 69.51A.040. A qualifying patient or designated provider meeting the conditions of this subsection but possessing more cannabis than the limits set forth in RCW 69.51A.040(1) may, in the investigating peace officer’s discretion, be taken into custody and booked into jail in connection with the investigation of the incident. [2011 c 181 § 402.]

*Reviser’s note: Section 901 of this act was vetoed by the governor.

69.51A.045 Possession of cannabis exceeding lawful amount—Affirmative defense. A qualifying patient or designated provider in possession of cannabis plants, useable cannabis, or cannabis product exceeding the limits set forth in RCW 69.51A.040(1) but otherwise in compliance with all other terms and conditions of this chapter may establish an affirmative defense to charges of violations of state law relating to cannabis through proof at trial, by a preponderance of the evidence, that he or she was a validly authorized qualifying patient or designated provider at the time of the officer’s questioning. A qualifying patient or designated provider who establishes an affirmative defense under the terms of this section may also establish an affirmative defense under RCW 69.51A.045. [2011 c 181 § 406.]

*Reviser’s note: The section creating a registry, 2011 c 181 § 901, was vetoed by the governor.

69.51A.050 Medical marijuana, lawful possession—State not liable. (1) The lawful possession or manufacture of medical marijuana as authorized by this chapter shall not result in the forfeiture or seizure of any property.

(2) No person shall be prosecuted for constructive possession, conspiracy, or any other criminal offense solely for being in the presence or vicinity of medical marijuana or its use as authorized by this chapter.

(3) The state shall not be held liable for any deleterious outcomes from the medical use of marijuana by any qualifying patient. [1999 c 2 § 7 (Initiative Measure No. 692, approved November 3, 1998).]

69.51A.055 Limitations of chapter—Persons under supervision. (1)(a) The arrest and prosecution protections established in RCW 69.51A.040 may not be asserted in a supervision revocation or violation hearing by a person who is supervised by a corrections agency or department, including local governments or jails, that has determined that the terms of this section are inconsistent with and contrary to his or her supervision.

(b) The affirmative defenses established in RCW 69.51A.043, 69.51A.045, 69.51A.047, and *section 407 of this act may not be asserted in a supervision revocation or violation hearing by a person who is supervised by a corrections agency or department, including local governments or jails, that has determined that the terms of this section are inconsistent with and contrary to his or her supervision.

(2) The provisions of RCW 69.51A.040, 69.51A.085, and 69.51A.025 do not apply to a person who is supervised for a criminal conviction by a corrections agency or department, including local governments or jails, that has determined that the terms of this chapter are inconsistent with and contrary to his or her supervision.

(3) A person may not be licensed as a licensed producer, licensed processor of cannabis products, or a licensed dispensary under *section 601, 602, or 701 of this act if he or she is supervised for a criminal conviction by a corrections agency or department, including local governments or jails, that has determined that licensure is inconsistent with and contrary to his or her supervision. [2011 c 181 § 1105.]

*Reviser’s note: Sections 407, 601, 602, and 701 were vetoed by the governor.

69.51A.060 Crimes—Limitations of chapter. (1) It shall be a class 3 civil infraction to use or display medical cannabis in a manner or place which is open to the view of the general public.

[Title 69 RCW—page 100]
(2) Nothing in this chapter establishes a right of care as a covered benefit or requires any state purchased health care as defined in RCW 41.05.011 or other health carrier or health plan as defined in Title 48 RCW to be liable for any claim for reimbursement for the medical use of cannabis. Such entities may enact coverage or noncoverage criteria or related policies for payment or nonpayment of medical cannabis in their sole discretion.

(3) Nothing in this chapter requires any health care professional to authorize the medical use of cannabis for a patient.

(4) Nothing in this chapter requires any accommodation of any on-site medical use of cannabis in any place of employment, in any school bus or on any school grounds, in any youth center, in any correctional facility, or smoking cannabis in any public place or hotel or motel.

(5) Nothing in this chapter authorizes the use of medical cannabis by any person who is subject to the Washington code of military justice in chapter 38.38 RCW.

(6) Employers may establish drug-free work policies. Nothing in this chapter requires an accommodation for the medical use of cannabis if an employer has a drug-free work place.

(7) It is a class C felony to fraudulently produce any record purporting to be, or tamper with the content of any record for the purpose of having it accepted as, valid documentation under *RCW 69.51A.010(32)(a), or to backdate such documentation to a time earlier than its actual date of execution.

(8) No person shall be entitled to claim the protection from arrest and prosecution under RCW 69.51A.040 or the affirmative defense under RCW 69.51A.043 for engaging in the medical use of cannabis in a way that endangers the health or well-being of any person through the use of a motorized vehicle on a street, road, or highway, including violations of RCW 46.61.502 or 46.61.504, or equivalent local ordinances. [2011 c 181 § 501; 2010 c 284 § 4; 2007 c 371 § 6; 1999 c 2 § 8 (Initiative Measure No. 692, approved November 3, 1998).]

*Reviser’s note: RCW 69.51A.010(32) is a reference to the definition of "valid documentation" which was amended and renumbered by 2011 c 181 § 201, but the section was vetoed by the governor.

*Reviser’s note: The section creating a registry, 2011 c 181 § 901, was vetoed by the governor.

69.51A.085 Collective gardens. (1) Qualifying patients may create and participate in collective gardens for the purpose of producing, processing, transporting, and delivering cannabis for medical use subject to the following conditions:

(a) No more than ten qualifying patients may participate in a single collective garden at any time;

(b) A collective garden may contain no more than fifteen plants per patient up to a total of forty-five plants;

(c) A collective garden may contain no more than twenty-four ounces of useable cannabis per patient up to a total of seventy-two ounces of useable cannabis;

(d) A copy of each qualifying patient’s valid documentation or proof of registration with the registry established in *section 901 of this act, including a copy of the patient’s proof of identity, must be available at all times on the premises of the collective garden; and

(e) No useable cannabis from the collective garden is delivered to anyone other than one of the qualifying patients participating in the collective garden.

(2) For purposes of this section, the creation of a "collective garden" means qualifying patients sharing responsibility for acquiring and supplying the resources required to produce and process cannabis for medical use such as, for example, a location for a collective garden; equipment, supplies, and labor necessary to plant, grow, and harvest cannabis; cannabis plants, seeds, and cuttings; and equipment, supplies, and labor necessary for proper construction, plumbing, wiring, and ventilation of a garden of cannabis plants.

(3) A person who knowingly violates a provision of subsection (1) of this section is not entitled to the protections of this chapter. [2011 c 181 § 403.]

69.51A.090 Applicability of valid documentation definition. The provisions of RCW 69.51A.010, relating to the definition of "valid documentation," apply prospectively only, not retroactively, and do not affect valid documentation obtained prior to June 10, 2010. [2010 c 284 § 5.]

69.51A.100 Qualifying patient’s designation of provider—Provider’s service as designated provider—Termination. (1) A qualifying patient may revoke his or her designation of a specific provider and designate a different provider at any time. A revocation of designation must be in writing, signed and dated. The protections of this chapter cease to apply to a person who has served as a designated provider to a qualifying patient seventy-two hours after receipt of that patient’s revocation of his or her designation.

(2) A person may stop serving as a designated provider to a given qualifying patient at any time. However, that person may not begin serving as a designated provider to a different qualifying patient until fifteen days have elapsed from the date the last qualifying patient designated him or her to serve as a provider. [2011 c 181 § 404.]
69.51A.110 Suitability for organ transplant. A qualifying patient’s medical use of cannabis as authorized by a health care professional may not be a sole disqualifying factor in determining the patient’s suitability for an organ transplant, unless it is shown that this use poses a significant risk of rejection or organ failure. This section does not preclude a health care professional from requiring that a patient abstain from the medical use of cannabis, for a period of time determined by the health care professional, while waiting for a transplant organ or before the patient undergoes an organ transplant. [2011 c 181 § 408.]

69.51A.120 Parental rights or residential time—Not to be restricted. A qualifying patient or designated provider may not have his or her parental rights or residential time with a child restricted solely due to his or her medical use of cannabis in compliance with the terms of this chapter absent written findings supported by evidence that such use has resulted in a long-term impairment that interferes with the performance of parenting functions as defined under RCW 26.09.004. [2011 c 181 § 409.]

69.51A.130 State and municipalities—Not subject to liability. (1) No civil or criminal liability may be imposed by any court on the state or its officers and employees for actions taken in good faith under this chapter and within the scope of their assigned duties.

(2) No civil or criminal liability may be imposed by any court on cities, towns, and counties or other municipalities and their officers and employees for actions taken in good faith under this chapter and within the scope of their assigned duties. [2011 c 181 § 1101.]

69.51A.140 Counties, cities, towns—Authority to adopt and enforce requirements. (1) Cities and towns may adopt and enforce any of the following pertaining to the production, processing, or dispensing of cannabis or cannabis products within their jurisdiction: Zoning requirements, business licensing requirements, health and safety requirements, and business taxes. Nothing in chapter 181, Laws of 2011 is intended to limit the authority of cities and towns to impose zoning requirements or other conditions upon licensed dispensers, so long as such requirements do not preclude the possibility of siting licensed dispensers within the jurisdiction. If the jurisdiction has no commercial zones, the jurisdiction is not required to adopt zoning to accommodate licensed dispensers.

(2) Counties may adopt and enforce any of the following pertaining to the production, processing, or dispensing of cannabis or cannabis products within their jurisdiction in locations outside of the corporate limits of any city or town: Zoning requirements, business licensing requirements, and health and safety requirements. Nothing in chapter 181, Laws of 2011 is intended to limit the authority of counties to impose zoning requirements or other conditions upon licensed dispensers, so long as such requirements do not preclude the possibility of siting licensed dispensers within the jurisdiction. If the jurisdiction has no commercial zones, the jurisdiction is not required to adopt zoning to accommodate licensed dispensers. [2011 c 181 § 1102.]

69.51A.200 Evaluation. (1) By July 1, 2014, the Washington state institute for public policy shall, within available funds, conduct a cost-benefit evaluation of the implementation of chapter 181, Laws of 2011 and the rules adopted to carry out its purposes.

(2) The evaluation of the implementation of chapter 181, Laws of 2011 and the rules adopted to carry out its purposes shall include, but not necessarily be limited to, consideration of the following factors:

(a) Qualifying patients’ access to an adequate source of cannabis for medical use;
(b) Qualifying patients’ access to a safe source of cannabis for medical use;
(c) Qualifying patients’ access to a consistent source of cannabis for medical use;
(d) Qualifying patients’ access to a secure source of cannabis for medical use;
(e) Qualifying patients’ and designated providers’ contact with law enforcement and involvement in the criminal justice system;
(f) Diversion of cannabis intended for medical use to nonmedical uses;
(g) Incidents of home invasion burglaries, robberies, and other violent and property crimes associated with qualifying patients accessing cannabis for medical use;
(h) Whether there are health care professionals who make a disproportionately high amount of authorizations in comparison to the health care professional community at large;
(i) Whether there are indications of health care professionals in violation of RCW 69.51A.030; and
(j) Whether the health care professionals making authorizations reside in this state or out of this state.

(3) For purposes of facilitating this evaluation, the departments of health and agriculture will make available to the Washington state institute for public policy requested data, and any other data either department may consider relevant, from which all personally identifiable information has been redacted. [2011 c 181 § 1001.]

69.51A.900 Short title—1999 c 2. This chapter may be known and cited as the Washington state medical use of cannabis act. [2011 c 181 § 1106; 1999 c 2 § 1 (Initiative Measure No. 692, approved November 3, 1998).]

69.51A.901 Severability—1999 c 2. If any provision of this act or its application to any person or circumstance is held invalid, the remainder of the act or the application of the provision to other persons or circumstances is not affected. [1999 c 2 § 10 (Initiative Measure No. 692, approved November 3, 1998).]

69.51A.902 Captions not law—1999 c 2. Captions used in this chapter are not any part of the law. [1999 c 2 § 11 (Initiative Measure No. 692, approved November 3, 1998).]

69.51A.903 Severability—2011 c 181. If any provision of this act or the application thereof to any person or circumstance is held invalid, the invalidity does not affect other provisions or applications of the act that can be given effect
without the invalid provision or application, and to this end the provisions of this act are severable. [2011 c 181 § 1103.]

Chapter 69.52 RCW
IMITATION CONTROLLED SUBSTANCES

Sections
69.52.010 Legislative findings.
69.52.020 Definitions.
69.52.030 Violations—Exceptions.
69.52.040 Seizure of contraband.
69.52.045 Seizure at rental premises—Notification of landlord.
69.52.050 Injunctive action by attorney general authorized.
69.52.060 Injunctive or other legal action by manufacturer of controlled substances authorized.
69.52.070 Violations—Juvenile driving privileges.
69.52.090 Severability—1982 c 171.
69.52.091 Effective date—1982 c 171.

Drug nuisances—Injunctions: Chapter 7.43 RCW.

69.52.010 Legislative findings. The legislature finds that imitation controlled substances are being manufactured to imitate the appearance of the dosage units of controlled substances for sale to school age youths and others to facilitate the fraudulent sale of controlled substances. The legislature further finds that manufacturers are endeavoring to profit from the manufacture of these imitation controlled substances while avoiding liability by accurately labeling the containers or packaging which contain these imitation controlled substances. The close similarity of appearance between dosage units of imitation controlled substances and controlled substances is indicative of a deliberate and wilful attempt to profit by deception without regard to the tragic human consequences. The use of imitation controlled substances is responsible for a growing number of injuries and deaths, and the legislature hereby declares that this chapter is necessary for the protection and preservation of the public health and safety. [1982 c 171 § 2.]

69.52.020 Definitions. Unless the context clearly requires otherwise, the definitions in this section apply throughout this chapter.

(1) "Controlled substance" means a substance as that term is defined in chapter 69.50 RCW.

(2) "Distribute" means the actual or constructive transfer (or attempted transfer) or delivery or dispensing to another of an imitation controlled substance.

(3) "Imitation controlled substance" means a substance that is not a controlled substance, but which by appearance or representation would lead a reasonable person to believe that the substance is a controlled substance. Appearance includes, but is not limited to, color, shape, size, and markings of the dosage unit. Representation includes, but is not limited to, representations or factors of the following nature:
(a) Statements made by an owner or by anyone else in control of the substance concerning the nature of the substance, or its use or effect;
(b) Statements made to the recipient that the substance may be resold for inordinate profit; or
(c) Whether the substance is packaged in a manner normally used for illicit controlled substances.

(4) "Manufacture" means the production, preparation, compounding, processing, encapsulating, packaging or repackaging, or labeling or relabeling of an imitation controlled substance. [1982 c 171 § 3.]

69.52.030 Violations—Exceptions. (1) It is unlawful for any person to manufacture, distribute, or possess with intent to distribute, an imitation controlled substance. Any person who violates this subsection shall, upon conviction, be guilty of a class C felony.

(2) Any person eighteen years of age or over who violates subsection (1) of this section by distributing an imitation controlled substance to a person under eighteen years of age is guilty of a class B felony.

(3) It is unlawful for any person to cause to be placed in any newspaper, magazine, handbill, or other publication, or to post or distribute in any public place, any advertisement or solicitation offering for sale imitation controlled substances. Any person who violates this subsection is guilty of a class C felony.

(4) No civil or criminal liability shall be imposed by virtue of this chapter on any person registered under the Uniform Controlled Substances Act pursuant to RCW 69.50.301 or 69.50.303 who manufactures, distributes, or possesses an imitation controlled substance for use as a placebo or other use by a registered practitioner, as defined in *RCW 69.50.101(t), in the course of professional practice or research.

(5) No prosecution under this chapter shall be dismissed solely by reason of the fact that the dosage units were contained in a bottle or other container with a label accurately describing the ingredients of the imitation controlled substance dosage units. The good faith of the defendant shall be an issue of fact for the trier of fact. [1983 1st ex.s. c 4 § 5; 1982 c 171 § 4.]

*Reviser’s note: The reference to RCW 69.50.101(t) is erroneous. "Practitioner" is defined in (w) of that section. RCW 69.50.101 was alphabetized pursuant to RCW 1.08.015(2)(k), changing subsection (w) to subsection (x).

Additional notes found at www.leg.wa.gov

69.52.040 Seizure of contraband. Imitation controlled substances shall be subject to seizure, forfeiture, and disposition in the same manner as are controlled substances under RCW 69.50.505. [1982 c 171 § 5.]

69.52.045 Seizure at rental premises—Notification of landlord. Whenever an imitation controlled substance which is manufactured, distributed, or possessed in violation of this chapter is seized at rental premises, the law enforcement agency shall make a reasonable attempt to discover the identity of the landlord and shall notify the landlord in writing, at the last address listed in the property tax records and at any other address known to the law enforcement agency, of the seizure and the location of the seizure. [1988 c 150 § 10.]

Legislative findings—Severability—1988 c 150: See notes following RCW 59.18.130.

69.52.050 Injunctive action by attorney general authorized. The attorney general is authorized to apply for injunctive action against a manufacturer or distributor of imitation controlled substances in this state. [1982 c 171 § 6.]

[Title 69 RCW—page 103]
69.52.060  Injunctive or other legal action by manufacturer of controlled substances authorized.  Any manufacturer of controlled substances licensed or registered in a state requiring such licensure or registration, may bring injunctive or other action against a manufacturer or distributor of imitation controlled substances in this state.  [1982 c 171 § 7.]

69.52.070  Violations—Juvenile driving privileges.  
(1) If a juvenile thirteen years of age or older and under the age of twenty-one is found by a court to have committed any offense that is a violation of this chapter, the court shall notify the department of licensing within twenty-four hours after entry of the judgment.  
(2) Except as otherwise provided in subsection (3) of this section, upon petition of a juvenile whose privilege to drive has been revoked pursuant to RCW 46.20.265, the court may at any time the court deems appropriate notify the department of licensing to reinstate the juvenile’s privilege to drive.  
(3) If the conviction is for the juvenile’s first violation of this chapter or chapter 66.44, 69.41, or 69.50 RCW, the juvenile may not petition the court for reinstatement of the juvenile’s privilege to drive revoked pursuant to RCW 46.20.265 until the later of ninety days after the date the juvenile turns sixteen or ninety days after the judgment was entered.  If the conviction was for the juvenile’s second or subsequent violation of this chapter or chapter 66.44, 69.41, or 69.50 RCW, the juvenile may not petition the court for reinstatement of the juvenile’s privilege to drive revoked pursuant to RCW 46.20.265 until the later of the date the juvenile turns seventeen or one year after the date judgment was entered.  [1989 c 271 § 121; 1988 c 148 § 6.]

Legislative finding—Severability—1988 c 148:  See notes following RCW 13.40.265.

Additional notes found at www.leg.wa.gov

69.52.900  Severability—1982 c 171.  If any provision of this chapter or its application to any person or circumstance is held invalid, the remainder of the chapter or the application of the provision to other persons or circumstances is not affected.  [1982 c 171 § 8.]

69.52.901  Effective date—1982 c 171.  This act shall take effect on July 1, 1982.  [1982 c 171 § 10.]

Chapter 69.53 RCW  
USE OF BUILDINGS FOR UNLAWFUL DRUGS

Sections

69.53.010  Unlawful use of building for drug purposes—Liability of owner or manager—Penalty.
69.53.020  Unlawful fortification of building for drug purposes—Penalty.
69.53.030  Unlawful use of fortified building—Penalty.

69.53.010  Unlawful use of building for drug purposes—Liability of owner or manager—Penalty.  (1) It is unlawful for any person who has under his or her management or control any building, room, space, or enclosure, either as an owner, lessee, agent, employee, or mortgagee, to knowingly rent, lease, or make available for use, with or without compensation, the building, room, space, or enclosure for the purpose of unlawfully manufacturing, delivering, selling, storing, or giving away any controlled substance under chapter 69.50 RCW, legend drug under chapter 69.41 RCW, or imitation controlled substance under chapter 69.52 RCW.

(2) It shall be a defense for an owner, manager, or other person in control pursuant to subsection (1) of this section to, in good faith, notify a law enforcement agency of suspected drug activity pursuant to subsection (1) of this section, or to process an unlawful detainer action for drug-related activity against the tenant or occupant.

(3) A violation of this section is a class C felony punishable under chapter 9A.20 RCW.  [1988 c 150 § 13; 1987 c 458 § 7.]

Legislative findings—Severability—1988 c 150:  See notes following RCW 59.18.130.

Additional notes found at www.leg.wa.gov

69.53.020  Unlawful fortification of building for drug purposes—Penalty.  (1) It is unlawful for any person who has under his or her management or control any building, room, space, or enclosure specifically designed to suppress law enforcement entry in order to further the unlawful manufacture, delivery, sale, storage, or gift of any controlled substance under chapter 69.50 RCW, legend drug under chapter 69.41 RCW, or imitation controlled substance under chapter 69.52 RCW.

(2) It shall be a defense for an owner, manager, or other person in control pursuant to subsection (1) of this section to, in good faith, notify a law enforcement agency of suspected drug activity pursuant to subsection (1) of this section, or to process an unlawful detainer action for drug-related activity against the tenant or occupant.

(3) A violation of this section is a class C felony punishable under chapter 9A.20 RCW.  [1988 c 150 § 14; 1987 c 458 § 8.]

Legislative findings—Severability—1988 c 150:  See notes following RCW 59.18.130.

Additional notes found at www.leg.wa.gov

69.53.030  Unlawful use of fortified building—Penalty.  (1) It is unlawful for any person to use a building, room, space, or enclosure specifically designed to suppress law enforcement entry in order to unlawfully manufacture, deliver, sell, store, or give away any controlled substance under chapter 69.50 RCW, legend drug under chapter 69.41 RCW, or imitation controlled substance under chapter 69.52 RCW.

(2) A violation of this section is a class C felony punishable under chapter 9A.20 RCW.  [1987 c 458 § 9.]

Additional notes found at www.leg.wa.gov

Chapter 69.55 RCW  
AMMONIA

(Formerly:  Anhydrous ammonia)

Sections

69.55.010  Theft of ammonia.
69.55.020  Unlawful storage of ammonia.
69.55.030  Damages—Liability.
**69.60.010** Theft of ammonia.  (1) A person who, with intent to deprive the owner or owner’s agent, wrongfully obtains pressurized ammonia gas or pressurized ammonia gas solution, is guilty of theft of ammonia.

(2) Theft of ammonia is a class C felony. [2002 c 133 § 1; 2000 c 225 § 1.]

**Effective date—2002 c 133:** "This act is necessary for the immediate preservation of the public peace, health, or safety, or support of the state government and its existing public institutions, and takes effect immediately [March 26, 2002]." [2002 c 133 § 5.]

Additional notes found at www.leg.wa.gov

**69.60.020** Unlawful storage of ammonia.  A person is guilty of the crime of unlawful storage of ammonia if the person possesses, transports, or delivers pressurized ammonia gas or pressurized ammonia gas solution in a container that (1) is not approved by the United States department of transportation to hold ammonia, or (2) was not constructed to meet state and federal industrial health and safety standards for holding ammonia. Violation of this section is a class C felony.

This section does not apply to public employees or private contractors authorized to clean up and dispose of hazardous waste or toxic substances under chapter 70.105 or 70.105D RCW or to solid waste haulers and their employees who unknowingly possess, transport, or deliver pressurized ammonia gas or pressurized ammonia gas solution during the course of the performance of their duties. [2002 c 133 § 2; 2000 c 225 § 2.]

**Effective date—2002 c 133:** See note following RCW 69.55.010.

Additional notes found at www.leg.wa.gov

**69.60.030** Damages—Liability.  Any damages arising out of the unlawful possession of, storage of, or tampering with pressurized ammonia gas or pressurized ammonia gas solution, or pressurized ammonia gas equipment or pressurized ammonia gas solution equipment, shall be the sole responsibility of the unlawful possessor, storer, or tamperer. In no case shall liability for damages arising out of the unlawful possession of, storage of, or tampering with pressurized ammonia gas or pressurized ammonia gas solution, or pressurized ammonia gas equipment or pressurized ammonia gas solution equipment, extend to the lawful owner, installer, maintainer, designer, manufacturer, possessor, or seller of the pressurized ammonia gas or pressurized ammonia gas solution, or pressurized ammonia gas equipment or pressurized ammonia gas solution equipment, unless such damages arise out of the owner, installer, maintainer, designer, manufacturer, possessor, or seller’s acts or omissions that constitute negligent misconduct to abide by the laws regarding pressurized ammonia gas or pressurized ammonia gas solution possession and storage. [2002 c 133 § 3; 2000 c 225 § 3.]

**Effective date—2002 c 133:** See note following RCW 69.55.010.

Additional notes found at www.leg.wa.gov

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**Chapter 69.60 RCW OVER-THE-COUNTER MEDICATIONS**

Sections

69.60.010 Legislative findings.
69.60.020 Definitions.
69.60.030 Identification required.

69.60.040 Imprint information—Publication—Availability.
69.60.050 Noncompliance—Contraband—Fine.
69.60.060 Rules.
69.60.070 Imprinting requirements—Retailers and wholesalers.
69.60.080 Exemptions—Application by manufacturer.
69.60.090 Implementation of federal system—Termination of state system.
69.60.901 Severability—1993 c 135.

**69.60.010 Legislative findings.** The legislature of the state of Washington finds that:

(1) Accidental and purposeful ingestions of solid medication forms continue to be the most frequent cause of poisoning in our state;

(2) Modern treatment is dependent upon knowing the ingredients of the ingestant;

(3) The imprinting of identifying characteristics on all tablets, capsules, and caplets of prescription medication forms, both trade name products and generic products, has been extremely beneficial in our state and was accomplished at trivial cost to the manufacturers and consumers;

(4) Although over-the-counter medications usually constitute a lower order of risk to inestees, treatment after overdose is equally dependent upon knowing the ingredients involved, but there is no coding index uniformly used by this class of medication;

(5) Approximately seventy percent of over-the-counter medications in solid form already have some type of identifier imprinted on their surfaces;

(6) While particular efforts are being instituted to prevent recurrent tampering with over-the-counter medications, the added benefit of rapid and prompt identification of all possible contaminated products, including over-the-counter medications, would make for a significant improvement in planning for appropriate tracking and monitoring programs;

(7) At the same time, health care professionals serving the elderly find it especially advantageous to be able to identify and confirm the ingredients of their multiple medications, including over-the-counter products, as are often consumed by such patients;

(8) The legislature supports and encourages efforts that are being made to establish a national, legally enforceable system governing the imprinting of solid dosage form over-the-counter medications, which system is consistent with the requirements of this chapter. [1989 c 247 § 1.]

**69.60.020 Definitions.** The terms defined in this section shall have the meanings indicated when used in this chapter.

(1) "Solid dosage form" means capsules or tablets or similar over-the-counter medication products intended for administration and which could be ingested orally.

(2) "Over-the-counter medication" means a drug that can be obtained without a prescription and is not restricted to use by prescribing practitioners. For purposes of this chapter, over-the-counter medication does not include vitamins.

(3) "Board" means the state board of pharmacy.

(4) "Purveyor" means any corporation, person, or other entity that offers over-the-counter medications for wholesale, retail, or other type of sale. [1989 c 247 § 3.]
69.60.030 Identification required. (1) No over-the-counter medication in solid dosage form may be manufactured or commercially distributed within this state unless it has clearly marked or imprinted on it an individual symbol, number, company name, words, letters, marking, or national drug code number identifying the medication and the manufacturer or distributor of the medication: PROVIDED, HOWEVER, That an over-the-counter medication which has clearly marked or imprinted on it a distinctive logo, symbol, product name, letters, or other identifying mark, or which by its color, shape, or size together with a distinctive logo, symbol, product name, letters, or other mark is identifiable, shall be deemed in compliance with the provisions of this chapter.

(2) No manufacturer may sell any over-the-counter medication in solid dosage form contained within a bottle, vial, carton, or other container, or in any way affixed or appended to or enclosed within a package of any kind designed or intended for delivery in such container or package to an ultimate consumer within this state unless such container or package has clearly and permanently marked or imprinted on it an individual symbol, number, company name, words, letters, marking, or national drug code number identifying the medication and the manufacturer, packer, or distributor of the medication. [1993 c 135 § 1; 1989 c 247 § 2.]

69.60.040 Imprint information—Publication—Availability. Each manufacturer shall publish and provide to the board printed material which will identify each current imprint used by the manufacturer and the board shall be notified of any change. This information shall be provided by the board to all pharmacies licensed in the state of Washington, poison control centers, and hospital emergency rooms. [1989 c 247 § 4.]

69.60.050 Noncompliance—Contraband—Fine. (1) Any over-the-counter medication prepared or manufactured or offered for sale in violation of this chapter or implementing rules shall be contraband and subject to seizure, in the same manner as contraband legend drugs under RCW 69.41.060.

(2) A purveyor who fails to comply with this chapter after one notice of noncompliance by the board is subject to a one thousand dollar civil fine for each instance of noncompliance. [1989 c 247 § 5.]

69.60.060 Rules. The board shall have authority to promulgate rules for the enforcement and implementation of this chapter. [1989 c 247 § 6.]

69.60.070 Imprinting requirements—Retailers and wholesalers. All over-the-counter medications manufactured, received by, distributed to, or shipped to any retailer or wholesaler in this state after January 1, 1994, shall meet the requirements of this chapter. No over-the-counter medication may be sold to a consumer in this state after January 1, 1995, unless such over-the-counter medication complies with the imprinting requirements of this chapter. [1993 c 135 § 2; 1989 c 247 § 7.]

69.60.080 Exemptions—Application by manufacturer. The board, upon application of a manufacturer, may exempt an over-the-counter drug from the requirements of chapter 69.60 RCW on the grounds that imprinting is infeasible because of size, texture, or other unique characteristics. [1989 c 247 § 8.]

69.60.090 Implementation of federal system—Termination of state system. Before January 1, 1994, the board of pharmacy will consult with the state toxicologist to determine whether the federal government has established a legally enforceable system that is substantially equivalent to the requirements of this chapter that govern the imprinting of solid dosage form over-the-counter medication. To be substantially equivalent, the effective dates for implementation of the federal system for imprinting solid dosage form over-the-counter medication must be the same or earlier than the effective dates of implementation set out in the state system for imprinting solid dosage form over-the-counter medication. If the board determines that the federal system for imprinting solid dosage form over-the-counter medication is substantially equivalent to the state system for imprinting solid dosage form over-the-counter medication, this chapter will cease to exist on January 1, 1994. If the board determines that the federal system is substantially equivalent, except that the federal dates for implementation are later than the Washington state dates, this chapter will cease to exist when the federal system is implemented. [1993 c 135 § 3; 1989 c 247 § 9.]

69.60.090 Severability—1993 c 135. If any provision of this act or its application to any person or circumstance is held invalid, the remainder of the act or the application of the provision to other persons or circumstances is not affected. [1993 c 135 § 4.]

69.60.091 Effective date—1993 c 135. This act is necessary for the immediate preservation of the public peace, health, or safety, or support of the state government and its existing public institutions, and shall take effect immediately [April 30, 1993]. [1993 c 135 § 5.]
69.80.020 Definitions. Unless the context clearly requires otherwise, the definitions in this section apply throughout this chapter.

(1) "Distributing organization" means a charitable nonprofit organization under section 501(c) of the federal internal revenue code which distributes food free of charge and includes any nonprofit organization that distributes food free of charge to other nonprofit organizations or to the public.

(2) "Donor" means a person, corporation, association, or other organization which donates food to a distributing organization. "Donor" includes, but is not limited to, farmers, processors, distributors, wholesalers, and retailers of food. "Donor" also includes persons who harvest agricultural crops or perishable foods which have been donated by the owner to a distributing organization.

(3) "Food" means food products for human consumption as defined in RCW 69.04.008. [1983 c 241 § 2.]

69.80.031 Good samaritan food donation act—Definitions—Collecting, distributing, gleaning—Liability. (1) This section may be cited as the "good samaritan food donation act."

(2) As used in this section:

(a) "Apparently fit grocery product" means a grocery product that meets all quality and labeling standards imposed by federal, state, and local laws and regulations even though the product may not be readily marketable due to appearance, age, freshness, grade, size, surplus, or other conditions.

(b) "Apparently wholesome food" means food that meets all quality and labeling standards imposed by federal, state, and local laws and regulations even though the food may not be readily marketable due to appearance, age, freshness, grade, size, surplus, or other conditions.

(c) "Donate" means to give without requiring anything of monetary value from the recipient, except that the term shall include giving by a nonprofit organization to another nonprofit organization, notwithstanding that the donor organization has charged a nominal fee to the donee organization, if the ultimate recipient or user is not required to give anything of monetary value.

(d) "Food" means a raw, cooked, processed, or prepared edible substance, ice, beverage, or ingredient used or intended for use in whole or in part for human consumption.

(e) "Gleaner" means a person who harvests for free distribution to the needy, or for donation to a nonprofit organization for ultimate distribution to the needy, an agricultural crop that has been donated by the owner.

(f) "Grocery product" means a nonfood grocery product, including a disposable paper or plastic product, household cleaning product, laundry detergent, cleaning product, or miscellaneous household item.

(g) "Gross negligence" means voluntary and conscious conduct by a person with knowledge, at the time of the conduct, that the conduct is likely to be harmful to the health or well-being of another person.

(h) "Intentional misconduct" means conduct by a person with knowledge, at the time of the conduct, that the conduct is harmful to the health or well-being of another person.

(i) "Nonprofit organization" means an incorporated or unincorporated entity that:

(i) Is operating for religious, charitable, or educational purposes; and

(ii) Does not provide net earnings to, or operate in any other manner that inures to the benefit of, any officer, employee, or shareholder of the entity.

(j) "Person" means an individual, corporation, partnership, organization, association, or governmental entity, including a retail grocer, wholesaler, hotel, motel, manufacturer, restaurant, caterer, farmer, and nonprofit food distributor or hospital. In the case of a corporation, partnership, organization, association, or governmental entity, the term includes an officer, director, partner, deacon, trustee, council member, or other elected or appointed individual responsible for the governance of the entity.

(3) A person or gleaner is not subject to civil or criminal liability arising from the nature, age, packaging, or condition of apparently wholesome food or an apparently fit grocery product that the person or gleaner donates in good faith to a nonprofit organization for ultimate distribution to needy individuals, except that this subsection does not apply to an injury to or death of an ultimate user or recipient of the food or grocery product that results from an act or omission of the donor constituting gross negligence or intentional misconduct.

(4) A person who allows the collection or gleaning of donations on property owned or occupied by the person by gleaners, or paid or unpaid representatives of a nonprofit organization, for ultimate distribution to needy individuals is not subject to civil or criminal liability that arises due to the injury or death of the gleaner or representative, except that this subsection does not apply to an injury or death that results from an act or omission of the person constituting gross negligence or intentional misconduct.

(5) If some or all of the donated food and grocery products do not meet all quality and labeling standards imposed by federal, state, and local laws and regulations, the person or gleaner who donates the food and grocery products is not subject to civil or criminal liability in accordance with this section if the nonprofit organization that receives the donated food or grocery products:

(a) Is informed by the donor of the distressed or defective condition of the donated food or grocery products;

(b) Agrees to recondition the donated food or grocery products to comply with all the quality and labeling standards prior to distribution; and

(c) Is knowledgeable of the standards to properly recondition the donated food or grocery product.

(6) This section may not be construed to create liability. [1994 c 299 § 36.]

Intent—Finding—Severability—Conflict with federal requirements—1994 c 299: See notes following RCW 74.12.400.

69.80.040 Information and referral service for food donation program. The department of agriculture shall maintain an information and referral service for persons and organizations that have notified the department of their desire to participate in the food donation program under this chapter. [1983 c 241 § 4.]

69.80.050 Inspection of donated food by state and local agencies—Variance. (1) Appropriate state and local
agencies are authorized to inspect donated food items for wholesomeness and may establish procedures for the handling of food items.

(2) To facilitate the free distribution of food to needy persons, the local health officer, upon request from either a donor or distributing organization, may grant a variance to chapter 246-215 WAC covering physical facilities, equipment standards, and food source requirements when no known or expected health hazard would exist as a result of the action. [2002 c 217 § 3; 1983 c 241 § 6.]

Effective date—2002 c 217 § 3: “Section 3 of this act is necessary for the immediate preservation of the public peace, health, or safety, or support of the state government and its existing public institutions, and takes effect immediately [March 28, 2002].” [2002 c 217 § 4.]

Finding—Purpose—2002 c 217: “The legislature finds and declares that the distribution of food by donors to charitable organizations, such as shelters, churches, and fraternal organizations, serving communal meals to needy individuals can be done safely consistent with rules and recommended health and safety guidelines. The establishment of recommended donor guidelines by the department of health can educate the public about the preparation and handling of food donated to charitable organizations for distribution to homeless and other needy people. The purpose of this act is to authorize and facilitate the donation of food to needy persons in accordance with health and safety guidelines and rules, to assure that the donated food will not place needy recipients at risk, and to encourage businesses and individuals to donate surplus food to charitable organizations serving our state’s needy population.” [2002 c 217 § 1.]

69.90.010 Definitions. Unless the context clearly requires otherwise, the definitions in this section apply throughout this chapter.

(1) "Food product" includes any article other than drugs, whether in raw or prepared form, liquid or solid, or packaged or unpackaged, and which is used for human consumption.

(2) "Kosher" means a food product which has been prepared, processed, manufactured, maintained, and sold in accordance with the requisites of traditional Jewish dietary law.

(3) "Person" includes individuals, partnerships, corporations, and associations. [1985 c 127 § 2.]

69.90.020 Sale of "kosher" and "kosher style" food products prohibited if not kosher—Representations—Penalty. (1) No person may knowingly sell or offer for sale any food product represented as "kosher" or "kosher style" when that person knows that the food product is not kosher and when the representation is likely to cause a prospective purchaser to believe that it is kosher. Such a representation can be made orally or in writing, or by display of a sign, mark, insignia, or simulation.

(2) A person violating this section is guilty of a gross misdemeanor. [2003 c 53 § 349; 1985 c 127 § 3.]

Intent—Effective date—2003 c 53: See notes following RCW 2.48.180.

69.90.030 Violation of chapter is violation of consumer protection act. A violation of this chapter shall constitute a violation of the consumer protection act, chapter 19.86 RCW. [1985 c 127 § 4.]

69.90.900 Short title. This chapter shall be known as the sale of kosher food products act of 1985. [1985 c 127 § 1.]