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LEGISLATIVE BILL 273

Introduced by Schrock, 38; Burling, 33; Chambers, 11; Cunningham, 18; Dierks, 40; Schimek, 27; Vrtiska, 1

Read first time January 5, 2001

Committee: Agriculture

A BILL

FOR AN ACT relating to agriculture; to amend sections 2-954, 16-230, and 17-563, Reissue Revised Statutes of Nebraska, and sections 28-401, 28-405, and 81-2,147.06, Revised Statutes Supplement, 2000; to provide for cultivation of industrial hemp; to harmonize provisions; and to repeal the original sections.

Be it enacted by the people of the State of Nebraska,
Section 1. Industrial hemp (cannabis sativa) having no more than three-tenths of one percent tetrahydrocannabinol is recognized as an oilseed. Upon meeting the requirements of section 2 of this act, any person in this state may plant, grow, harvest, possess, process, sell, and buy industrial hemp (cannabis sativa) having no more than three-tenths of one percent tetrahydrocannabinol.

Sec. 2. (1) Any person desiring to grow industrial hemp for commercial purposes shall apply to the Department of Agriculture for a license on a form prescribed by the department. The application for a license shall include the name and address of the applicant and the legal description of the land area to be used for the production of industrial hemp. Except for employees of the agricultural experiment station or the Cooperative Extension Service of the University of Nebraska involved in research and extension-related activities, each applicant for initial licensure shall file a set of the applicant's fingerprints and any other information necessary to complete a check of his or her criminal history record information maintained by the Identification Division of the Federal Bureau of Investigation through the Nebraska State Patrol. All costs associated with such check are the responsibility of the applicant. Criminal history records provided to the department under this section are confidential. The department may use the records only in determining an applicant's eligibility for licensure under this section. Any person with a prior criminal conviction is not eligible for licensure under this section. If the applicant has completed the application process to the satisfaction of the department, the
department shall issue the license which shall be valid for a period of one year. Any person licensed under this section is presumed to be growing industrial hemp for commercial purposes.

Sec. 3. Each industrial hemp licensee shall file with the Department of Agriculture documentation indicating that the industrial seeds planted were of a type and variety certified to have no more than three-tenths of one percent tetrahydrocannabinol and a copy of any contract to grow industrial hemp. Each licensee shall notify the department of the sale or distribution of any industrial hemp grown by the licensee and the names of the persons to whom the hemp was sold or distributed.

Sec. 4. The Department of Agriculture shall adopt and promulgate rules and regulations to allow the industrial hemp to be tested during growth for tetrahydrocannabinol levels and to allow for supervision of the industrial hemp during its growth and harvest. To provide sufficient funds to pay costs associated with monitoring and testing industrial hemp in the state, the department shall assess each industrial hemp licensee a fee of five dollars per acre. The minimum fee assessed shall be one hundred fifty dollars per licensee. Such fees shall be remitted to the State Treasurer for credit to the Industrial Hemp Licensure Fund, which is hereby created. Money in the fund shall be used by the department to carry out and enforce sections 1 to 4 of this act. Any money in the fund available for investment shall be invested by the state investment officer pursuant to the Nebraska Capital Expansion Act and the Nebraska State Funds Investment Act.

Sec. 5. Section 2-954, Reissue Revised Statutes of Nebraska, is amended to read:
2-954. (1)(a) The duty of enforcing and carrying out the Noxious Weed Control Act shall be vested in the director and the control authorities as designated in the act. The director shall determine what weeds are noxious for purposes of the act. Industrial hemp (cannabis sativa) having no more than three-tenths of one percent tetrahydrocannabinol shall not be designated as a noxious weed. A list of such noxious weeds shall be included in the rules and regulations adopted and promulgated by the director.

The director shall prepare, publish, and revise as necessary a list of noxious weeds. The list shall be distributed to the public by the director, the Cooperative Extension Service, the control authorities, and any other body the director deems appropriate. The director shall, from time to time, adopt and promulgate rules and regulations on methods for control of noxious weeds and adopt and promulgate such rules and regulations as are necessary to carry out the act. Whenever special weed control problems exist in a county involving weeds not included in the rules and regulations, the control authority may petition the director to bring such weeds under the county control program. The petition shall contain the approval of the county board. Prior to petitioning the director, the control authority, in cooperation with the county board, shall hold a public hearing and take testimony upon the petition. Such hearing and the notice thereof shall be in the manner prescribed by the Administrative Procedure Act. A copy of the transcript of the public hearing shall accompany the petition filed with the director. The director may approve or disapprove the request. If approval is granted, the control authority may proceed under the forced control provisions of sections 2-953 to 2-955 and 2-958.
(b) The director shall (i) investigate the subject of noxious weeds, (ii) require information and reports from any control authority as to the presence of noxious weeds and other information relative to noxious weeds and the control thereof in localities where such control authority has jurisdiction, (iii) cooperate with control authorities in carrying out other laws administered by him or her, (iv) cooperate with agencies of federal and state governments and other persons in carrying out his or her duties under the Noxious Weed Control Act, (v) with the consent of the Governor, conduct investigations outside this state to protect the interest of the agricultural industry of this state from noxious weeds not generally distributed therein, (vi) with the consent of the federal agency involved, control noxious weeds on federal lands within this state, with reimbursement, when deemed by the director to be necessary to an effective weed control program, (vii) advise and confer as to the extent of noxious weed infestations and the methods determined best suited to the control thereof, (viii) call and attend meetings and conferences dealing with the subject of noxious weeds, (ix) disseminate information and conduct educational campaigns with respect to control of noxious weeds, (x) procure materials and equipment and employ personnel necessary to carry out the director's duties and responsibilities, and (xi) perform such other acts as may be necessary or appropriate to the administration of the act.

(c) When the director determines that a control authority has substantively failed to carry out its duties and responsibilities as a control authority or has substantively failed to implement a county weed control program, he or she shall
instruct the control authority regarding the measures necessary to fulfill such duties and responsibilities. The director shall establish a reasonable date by which the control authority shall fulfill such duties and responsibilities. If the control authority fails or refuses to comply with instructions by such date, the Attorney General shall file an action as provided by law against the control authority for such failure or refusal.

(2)(a) Each control authority shall carry out the duties and responsibilities vested in it under the act with respect to land under its jurisdiction in accordance with rules and regulations adopted and promulgated by the director. Such duties shall include the establishment of a coordinated program for control of noxious weeds within the county.

(b) A control authority may cooperate with any person in carrying out its duties and responsibilities under the act.

(3)(a) Each county board shall employ one or more weed control superintendents. Each such superintendent shall, as a condition precedent to employment, be certified in writing by the federal Environmental Protection Agency as a commercial applicator under the Federal Insecticide, Fungicide, and Rodenticide Act. Each superintendent shall be bonded for such sum as the county board shall prescribe. The same person may be a weed control superintendent for more than one county. Such employment may be for such tenure and at such rates of compensation and reimbursement for travel expenses as the county board may prescribe. Such superintendent shall be reimbursed for mileage at a rate equal to or greater than the rate provided in section 81-1176.

(b) Under the direction of the control authority, it
shall be the duty of every weed control superintendent to examine all land under the jurisdiction of the control authority for the purpose of determining whether the Noxious Weed Control Act and the rules and regulations adopted and promulgated by the director have been complied with. The weed control superintendent shall: (i) Compile such data on infested areas and controlled areas and such other reports as the director or the control authority may require; (ii) consult and advise upon matters pertaining to the best and most practical methods of noxious weed control and render assistance and direction for the most effective control; (iii) investigate or aid in the investigation and prosecution of any violation of the act; and (iv) perform such other duties as required by the control authority in the performance of its duties. Weed control superintendents shall cooperate and assist one another to the extent practicable and shall supervise the carrying out of the coordinated control program within the county.

(c) In cases involving counties in which municipalities have ordinances for weed control, the control authority may enter into agreements with municipal authorities for the enforcement of local weed ordinances and may follow collection procedures established by such ordinances. All money received shall be deposited in the weed control authority fund.

Sec. 6. Section 16-230, Reissue Revised Statutes of Nebraska, is amended to read:

16-230. (1) A city of the first class by ordinance may require lots or pieces of ground within the city or within two miles of the corporate limits of the city to be drained or filled so as to prevent stagnant water or any other nuisance accumulating
thereon. It may require the owner or occupant of all lots and
pieces of ground within the city to keep the lots and pieces of
ground and the adjoining streets and alleys free of any growth of
twelve inches or more in height of weeds, grasses, or worthless
vegetation, and it may prohibit and control the throwing,
depositing, or accumulation of litter on any lot or piece of ground
within the city.

(2) Any city of the first class may by ordinance declare
it to be a nuisance to permit or maintain any growth of twelve
inches or more in height of weeds, grasses, or worthless vegetation
or to litter or cause litter to be deposited or remain thereon
except in proper receptacles.

(3) Any owner or occupant of a lot or piece of ground
shall, upon conviction of violating such ordinance, be guilty of a
Class V misdemeanor.

(4) Notice to abate and remove such nuisance shall be
given to each owner or owner's duly authorized agent and to the
occupant, if any, by personal service or certified mail. Within
five days after receipt of such notice, if the owner or occupant of
the lot or piece of ground does not request a hearing with the city
or fails to comply with the order to abate and remove the nuisance,
the city may have such work done. The costs and expenses of any
such work shall be paid by the owner. If unpaid for two months
after such work is done, the city may either (a) levy and assess
the costs and expenses of the work upon the lot or piece of ground
so benefited in the same manner as other special taxes for
improvements are levied and assessed or (b) recover in a civil
action the costs and expenses of the work upon the lot or piece of
ground and the adjoining streets and alleys.

(5) For purposes of this section:

(a) Litter shall include, but not be limited to: (i) Trash, rubbish, refuse, garbage, paper, rags, and ashes; (ii) wood, plaster, cement, brick, or stone building rubble; (iii) grass, leaves, and worthless vegetation; (iv) offal and dead animals; and (v) any machine or machines, vehicle or vehicles, or parts of a machine or vehicle which have lost their identity, character, utility, or serviceability as such through deterioration, dismantling, or the ravages of time, are inoperative or unable to perform their intended functions, or are cast off, discarded, or thrown away or left as waste, wreckage, or junk; and

(b) Weeds shall include, but not be limited to, bindweed (Convolvulus arvensis), puncture vine (Tribulus terrestris), leafy spurge (Euphorbia esula), Canada thistle (Cirsium arvense), perennial peppergrass (Lepidium draba), Russian knapweed (Centaurea picris), Johnson grass (Sorghum halepense), nodding or musk thistle, quack grass (Agropyron repens), perennial sow thistle (Sonchus arvensis), horse nettle (Solanum carolinense), bull thistle (Cirsium lanceolatum), buckthorn (Rhamnus sp.) (tourn), hemp plant (Cannabis sativa) having more than three-tenths of one percent tetrahydrocannabinol, and ragweed (Ambrosiaceae).

Sec. 7. Section 17-563, Reissue Revised Statutes of Nebraska, is amended to read:

17-563. (1) Each city of the second class and village by ordinance may require lots or pieces of ground within the city or village to be drained or filled so as to prevent stagnant water or any other nuisance accumulating thereon. It may require the owner
or occupant of any lot or piece of ground within the city or village to keep the lot or piece of ground and the adjoining streets and alleys free of any growth of twelve inches or more in height of weeds, grasses, or worthless vegetation, and it may prohibit and control the throwing, depositing, or accumulation of litter on any lot or piece of ground within the city or village.

(2) Any city of the second class and village may by ordinance declare it to be a nuisance to permit or maintain any growth of twelve inches or more in height of weeds, grasses, or worthless vegetation or to litter or cause litter to be deposited or remain thereon except in proper receptacles.

(3) Any owner or occupant of a lot or piece of ground shall, upon conviction of violating such ordinance, be guilty of a Class V misdemeanor.

(4) Notice to abate and remove such nuisance shall be given to each owner or owner’s duly authorized agent and to the occupant, if any, by personal service or certified mail. Within five days after receipt of such notice, if the owner or occupant of the lot or piece of ground does not request a hearing with the city or village or fails to comply with the order to abate and remove the nuisance, the city or village may have such work done. The costs and expenses of any such work shall be paid by the owner. If unpaid for two months after such work is done, the city or village may either (a) levy and assess the costs and expenses of the work upon the lot or piece of ground so benefited in the same manner as other special taxes for improvements are levied and assessed or (b) recover in a civil action the costs and expenses of the work upon the lot or piece of ground and the adjoining streets and alleys.
(5) For purposes of this section:

(a) Litter shall include, but not be limited to: (i) Trash, rubbish, refuse, garbage, paper, rags, and ashes; (ii) wood, plaster, cement, brick, or stone building rubble; (iii) grass, leaves, and worthless vegetation; (iv) offal and dead animals; and (v) any machine or machines, vehicle or vehicles, or parts of a machine or vehicle which have lost their identity, character, utility, or serviceability as such through deterioration, dismantling, or the ravages of time, are inoperative or unable to perform their intended functions, or are cast off, discarded, or thrown away or left as waste, wreckage, or junk; and

(b) Weeds shall include, but not be limited to, bindweed (Convolvulus arvensis), puncture vine (Tribulus terrestris), leafy spurge (Euphorbia esula), Canada thistle (Cirsium arvense), perennial peppergrass (Lepidium draba), Russian knapweed (Centaurea picris), Johnson grass (Sorghum halepense), nodding or musk thistle, quack grass (Agropyron repens), perennial sow thistle (Sonchus arvensis), horse nettle (Solanum carolinense), bull thistle (Cirsium lanceolatum), buckthorn (Rhamnus sp.) (tourn), hemp plant (Cannabis sativa) having more than three-tenths of one percent tetrahydrocannabinol, and ragweed (Ambrosiaceae).

Sec. 8. Section 28-401, Revised Statutes Supplement, 2000, is amended to read:

28-401. As used in the Uniform Controlled Substances Act, unless the context otherwise requires:

(1) Administer shall mean the direct application of a controlled substance, whether by injection, inhalation, ingestion, or any other means, to the body of a patient or research subject
by: (a) A practitioner or, in his or her presence, by his or her authorized agent; or (b) the patient or research subject at the direction and in the presence of the practitioner;

(2) Agent shall mean an authorized person who acts on behalf of or at the direction of a manufacturer, distributor, or dispenser. Agent shall not include a common or contract carrier, public warehouse keeper, or employee of the carrier or warehouse keeper;

(3) Administration shall mean the Drug Enforcement Administration, United States Department of Justice;

(4) Controlled substance shall mean a drug, substance, or immediate precursor in Schedules I to V of section 28-405. Controlled substance shall not include distilled spirits, wine, malt beverages, tobacco, or any nonnarcotic substance if such substance may, under the Federal Food, Drug, and Cosmetic Act and the law of this state, be lawfully sold over the counter without a prescription;

(5) Counterfeit substance shall mean 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 or persons who in fact manufactured, distributed, or dispensed such substance and which thereby falsely purports or is represented to be the product of, or to have been distributed by, such other manufacturer, distributor, or dispenser;

(6) Department shall mean the Department of Health and Human Services Regulation and Licensure personnel who are
responsible for the enforcement of the Uniform Controlled Substances Act in the areas assigned to it by the act;

(7) Division of Drug Control shall mean the personnel of the Nebraska State Patrol who are assigned to enforce the Uniform Controlled Substances Act;

(8) Dispense shall mean to deliver a controlled substance to an ultimate user or a research subject pursuant to the lawful order or prescription of a physician, physician assistant, dentist, veterinarian, or other medical practitioner licensed under the laws of this state to prescribe drugs, including the packaging, labeling, or compounding necessary to prepare the substance for such delivery. Dispenser shall mean the apothecary, pharmacist, or other practitioner, duly licensed, who dispenses a controlled substance to an ultimate user or a research subject;

(9) Distribute shall mean to deliver other than by administering or dispensing a controlled substance. Distributor shall mean a person who so distributes a controlled substance;

(10) Prescribe shall mean the act of a physician, physician assistant, surgeon, dentist, veterinarian, or other medical practitioner licensed under the laws of this state in issuing an order, prescription, or direction to a pharmacist or pharmacy to dispense a drug as required by the laws of this state;

(11) Drug shall mean (a) articles recognized in the official United States Pharmacopoeia, official Homeopathic Pharmacopoeia of the United States, 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, and (c) substances intended for
use as a component of any article specified in subdivision (a) or (b) of this subdivision, but shall not include devices or their components, parts, or accessories;

(12) Deliver or delivery shall mean the actual, constructive, or attempted transfer from one person to another of a controlled substance, whether or not there is an agency relationship;

(13) Marijuana shall mean all parts of the plant of the genus cannabis having more than three-tenths of one percent tetrahydrocannabinol, whether growing or not, the seeds thereof, and every compound, manufacture, salt, derivative, mixture, or preparation of such plant or its seeds, but shall not include the mature stalks of such plant, hashish, tetrahydrocannabinols extracted or isolated from the plant, fiber produced from such stalks, oil or cake made from the seeds of such plant, any other compound, manufacture, salt, derivative, mixture, or preparation of such mature stalks, or the sterilized seed of such plant which is incapable of germination. When the weight of marijuana is referred to in the Uniform Controlled Substances Act, it shall mean its weight at or about the time it is seized or otherwise comes into the possession of law enforcement authorities, whether cured or uncured at that time;

(14) Manufacture shall mean the production, preparation, propagation, compounding, or processing of a controlled substance, either directly or indirectly by extraction from substances of natural origin, independently by means of chemical synthesis, or by a combination of extraction and chemical synthesis, and shall include any packaging or repackaging of the substance or labeling
or relabeling of its container, except that manufacture shall not
include the preparation or compounding of a controlled substance by
an individual for his or her own use or the preparation,
compounding, packaging, or labeling of a controlled substance: (a)
By a practitioner as an incident to his or her prescribing,
administering, or dispensing of a controlled substance in the
course of his or her professional practice; or (b) by a
practitioner, or by his or her authorized agent under his or her
supervision, for the purpose of, or as an incident to, research,
teaching, or chemical analysis and not for sale;
(15) Narcotic drug shall mean any of the following,
whether produced directly or indirectly by extraction from
substances of vegetable origin, independently by means of chemical
synthesis, or by a combination of extraction and chemical
synthesis: (a) Opium, opium poppy and poppy straw, coca leaves, and
opiates; (b) a compound, manufacture, salt, derivative, or
preparation of opium, coca leaves, or opiates; or (c) a substance
and any compound, manufacture, salt, derivative, or preparation
thereof which is chemically equivalent to or identical with any of
the substances referred to in subdivisions (a) and (b) of this
subdivision, except that the words narcotic drug as used in the
Uniform Controlled Substances Act shall not include decocainized
coca leaves or extracts of coca leaves, which extracts do not
contain cocaine or ecgonine, or isoquinoline alkaloids of opium;
(16) Opiate shall mean any substance having an
addiction-forming or addiction-sustaining liability similar to
morphine or being capable of conversion into a drug having such
addiction-forming or addiction-sustaining liability. Opiate shall
not include the dextrorotatory isomer of 3-methoxy-n
methylnorphinane and its salts. Opiate shall include its racemic
and levorotatory forms;

(17) Opium poppy shall mean the plant of the species
Papaver somniferum L., except the seeds thereof;

(18) Poppy straw shall mean all parts, except the seeds,
of the opium poppy after mowing;

(19) Person shall mean any corporation, association,
partnership, limited liability company, or one or more individuals;

(20) Practitioner shall mean a physician, physician
assistant, dentist, veterinarian, pharmacist, scientific
investigator, pharmacy, or hospital, licensed, registered, or
otherwise permitted to distribute, dispense, prescribe, conduct
research with respect to, or administer a controlled substance in
the course of professional practice or research in this state, or
other person licensed, registered, or otherwise permitted to
distribute, dispense, conduct research with respect to, or
administer a controlled substance in the course of professional
practice or research in this state;

(21) Production shall include the manufacture, planting,
cultivation, or harvesting of a controlled substance;

(22) Immediate precursor shall mean a substance which is
the principal compound commonly used or produced primarily for use
and which is an immediate chemical intermediary used or likely to
be used in the manufacture of a controlled substance, the control
of which is necessary to prevent, curtail, or limit such
manufacture;

(23) State shall mean the State of Nebraska;
(24) Ultimate user shall mean a person who lawfully possesses a controlled substance for his or her own use, for the use of a member of his or her household, or for administration to an animal owned by him or her or by a member of his or her household;

(25) Physician shall mean a person authorized by law to practice medicine in this state and any other person authorized by law to treat sick and injured human beings in this state;

(26) Dentist shall mean a person authorized by law to practice dentistry in this state;

(27) Veterinarian shall mean a person authorized by law to practice veterinary medicine in this state;

(28) Hospital shall mean an institution for the care and treatment of sick and injured human beings and approved by the department;

(29) Podiatrist shall mean a person authorized by law to practice podiatry and who has graduated from an accredited school of podiatry in or since 1935;

(30) Apothecary shall mean a licensed pharmacist as defined by the laws of this state and, when the context so requires, the owner of the store or other place of business where drugs are compounded or dispensed by a licensed pharmacist, but nothing in this subdivision shall be construed as conferring on a person who is not registered nor licensed as a pharmacist any authority, right, or privilege that is not granted to him or her by the pharmacy laws of this state;

(31) Nothing in the Uniform Controlled Substances Act shall be construed as authority for a practitioner to perform an
act for which he or she is not authorized by the laws of this state;

(32) Cooperating individual shall mean any person, other than a commissioned law enforcement officer, who acts on behalf of, at the request of, or as agent for a law enforcement agency for the purpose of gathering or obtaining evidence of offenses punishable under the Uniform Controlled Substances Act;

(33) Hashish or concentrated cannabis shall mean: (a) The separated resin, whether crude or purified, obtained from a plant of the genus cannabis having more than three-tenths of one percent tetrahydrocannabinol; or (b) any material, preparation, mixture, compound, or other substance which contains ten percent or more by weight of tetrahydrocannabinols;

(34) Exceptionally hazardous drug shall mean (a) a narcotic drug, (b) thiophene analog of phencyclidine, (c) phencyclidine, (d) amobarbital, (e) secobarbital, or (f) pentobarbital;

(35) Imitation controlled substance shall mean a substance which is not a controlled substance but which, by way of express or implied representations and consideration of other relevant factors including those specified in section 28-445, would lead a reasonable person to believe the substance is a controlled substance. A placebo or registered investigational drug manufactured, distributed, possessed, or delivered in the ordinary course of practice or research by a health care professional shall not be deemed to be an imitation controlled substance;

(36) Controlled substance analogue shall mean a substance (a) the chemical structure of which is substantially similar to the
chemical structure of a Schedule I or Schedule II controlled substance as provided in section 28-405 or (b) which has a stimulant, depressant, analgesic, or hallucinogenic effect on the central nervous system that is substantially similar to or greater than the stimulant, depressant, analgesic, or hallucinogenic effect on the central nervous system of a Schedule I or Schedule II controlled substance as provided in section 28-405. A controlled substance analogue shall, to the extent intended for human consumption, be treated as a controlled substance under Schedule I of section 28-405 for purposes of the Uniform Controlled Substances Act. Controlled substance analogue shall not include (i) a controlled substance, (ii) any substance generally recognized as safe and effective within the meaning of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 301 et seq., (iii) any substance for which there is an approved new drug application, or (iv) with respect to a particular person, any substance if an exemption is in effect for investigational use for that person, under section 505 of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355, to the extent conduct with respect to such substance is pursuant to such exemption;

(37) Anabolic steroid shall mean any drug or hormonal substance, chemically and pharmacologically related to testosterone, (other than estrogens, progestins, and corticosteroids) that promotes muscle growth and includes any controlled substance in Schedule III(d) of section 28-405. Anabolic steroid shall not include any anabolic steroid which is expressly intended for administration through implants to cattle or other nonhuman species and has been approved by the Secretary of
Health and Human Services for such administration, but if any person prescribes, dispenses, or distributes such a steroid for human use, such person shall be considered to have prescribed, dispensed, or distributed an anabolic steroid within the meaning of this subdivision; and

(38) Physician assistant shall mean an individual licensed in accordance with sections 71-1,107.15 to 71-1,107.30.

Sec. 9. Section 28-405, Revised Statutes Supplement, 2000, is amended to read:

28-405. The following are the schedules of controlled substances referred to in the Uniform Controlled Substances Act:

Schedule I

(a) Any of the following opiates, including their isomers, esters, ethers, salts, and salts of isomers, esters, and ethers, unless specifically excepted, whenever the existence of such isomers, esters, ethers, and salts is possible within the specific chemical designation: (1) Acetylmethadol; (2) allylprodine; (3) alphacetylmethadol, except levo-alphacetylmethadol which is also known as levo-alpha-acetylmethadol, levomethadyl acetate, and LAAM; (4) alphameprodine; (5) alphamethadol; (6) benzethidine; (7) betacetylmethadol; (8) betameprodine; (9) betamethadol; (10) betaprodine; (11) clonitazene; (12) dextromoramide; (13) difenoxin; (14) diampromide; (15) diethylthiambutene; (16) dimenoxadol; (17) dimepheptanol; (18) dimethylthiambutene; (19) dioxaphetyl butyrate; (20) dipipanone; (21) ethylmethylthiambutene; (22) etonitazene; (23) etoxeridine; (24) furethidine; (25) hydroxypethidine; (26) ketobemidone; (27) levomoramide; (28) levophenacylmorphan; (29) -20-
1 morpheridine; (30) noracymethadol; (31) norlevorphanol; (32) normethadone; (33) norpipanone; (34) phenadoxone; (35) phenampronide; (36) phenomorphan; (37) phenoperidine; (38) piritramide; (39) proheptazine; (40) properidine; (41) propiram; (42) racemoramide; (43) trimeperidine; (44) alpha-methylfentanyl, N-(1-(alpha-methyl-beta-phenyl)ethyl-4-piperidyl) propionanilide, 1-(1-methyl-2-phenylethyl)-4-(N-propanilido) piperidine; (45) tilidine; (46) 3-Methylfentanyl, N-(3-methyl-1-(2-phenylethyl)-4-piperidyl)-N-phenylpropanamide, its optical and geometric isomers, salts, and salts of isomers; (47) 1-methyl-4-phenyl-4-propionoxypiperidine (MPPP), its optical isomers, salts, and salts of isomers; (48) 1-(2-phenylethyl)-4-phenyl-4-acetyloxy piperidine (PEPAP), its optical isomers, salts, and salts of isomers; (49) N-(1-(1-methyl-2-phenyl)ethyl-4-piperidyl)-N-phenylacetamide (acetyl-alpha-methylfentanyl), its optical isomers, salts, and salts of isomers; (50) N-(1-(1-methyl-2-(2-thienyl)ethyl-4-piperidyl)-Nphenylpropanamide (alpha-methylthiofentanyl), its optical isomers, salts, and salts of isomers; (51) N-(1-benxyl-4-piperidyl)-N-phenylpropanamide (benzylfentanyl), its optical isomers, salts, and salts of isomers; (52) N-(1-(2-hydroxy-2-phenyl)ethyl-4-piperidyl)-N-phenylpropanamide (beta-hydroxyfentanyl), its optical isomers, salts, and salts of isomers; (53) N-(3-methyl-1-(2-hydroxy-2-phenyl)ethyl-4-piperidyl)-N-phenylpropanamide (beta-hydroxy-3-methylfentanyl), its optical and geometric isomers, salts, and salts of isomers; (54) N-(3-methyl-1-(2-(2-thienyl)ethyl-4-piperidyl)-N-phenylpropanamide
(3-methylthiofentanyl), its optical and geometric isomers, salts, and salts of isomers; (55) N-(1-(2-thienyl)methyl-4-piperidyl)-N-phenylpropanamide (thenylfentanyl), its optical isomers, salts, and salts of isomers; (56) N-(1-(2-(2-thienyl)ethyl)-4-piperidyl)-N-phenylpropanamide (thiofentanyl), its optical isomers, salts, and salts of isomers; and (57) N-(1-(2-phenylethyl)-4-piperidyl)-N-(4-fluorophenyl)-propanamide (para-fluorofentanyl), its optical isomers, salts, and salts of isomers.

(b) Any of the following opium derivatives, their salts, isomers, and salts of isomers, unless specifically excepted, whenever the existence of such 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; and (23) thebacon.

(c) Any material, compound, mixture, or preparation which contains any quantity of the following hallucinogenic substances, their salts, isomers, and salts of isomers, unless specifically excepted, whenever the existence of such salts, isomers, and salts of isomers is possible within the specific chemical designation, and, for purposes of this subdivision only, isomer shall include the optical, position, and geometric isomers: (1) Bufotenine. Trade and other names shall include, but are not limited to:
3-[(B-Dimethylaminoethyl)-5-hydroxyindole;  
3-(2-dimethylaminoethyl)-5-indolol; N, N-dimethylserotonin;  
5-hydroxy-N, N-dimethyltryptamine; and mappine; (2)  
dimethyltryptamine. Trade and other names shall include, but are  
not limited to: N, N-diethyltryptamine; and DET; (3)  
dimethyltryptamine. Trade and other names shall include, but are  
not limited to: DMT; (4) 4-bromo-2, 5-dimethoxyamphetamine. Trade  
and other names shall include, but are not limited to: 4-bromo-2,  
5-dimethoxy-a-methylphenethylamine; and 4-bromo-2, 5-DMA; (5)  
4-methoxyamphetamine. Trade and other names shall include, but are  
not limited to: 4-methoxy-a-methyl-phenethylamine; and  
paramethoxyamphetamine, PMA; (6) 4-methyl-2,  
5-dimethoxyamphetamine. Trade and other names shall include, but  
are not limited to: 4-methyl-2, 5-dimethoxy-a-methylphenethylamine;  
DOM; and STP; (7) 5-methoxy-N-N, dimethyltryptamine; (8) ibogaine.  
Trade and other names shall include, but are not limited to:  
7-ethyl-6, 6B, 7, 8, 9, 10, 12, 13-octahydro-2-methoxy-6,  
9-methano-5H-pyrido (1',2':1,2) azepino (5, 4-b) indole; and  
tabernanthe iboga; (9) lysergic acid diethylamide; (10) marijuana  
having more than three-tenths of one percent tetrahydrocannabinol;  
(11) mescaline; (12) peyote. Peyote shall mean 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 or its  
seeds or extracts; (13) psilocybin; (14) psilocyn; (15)  
tetrahydrocannabinols of more than three-tenths of one percent,  
including, but not limited to, synthetic equivalents of the
substances contained in the plant or in the resinous extractives of
cannabis, sp. or synthetic substances, derivatives, and their
isomers with similar chemical structure and pharmacological
activity such as the following: Delta 1 cis or trans
tetrahydrocannabinol and their optical isomers, excluding
dronabinol in sesame oil and encapsulated in a soft gelatin capsule
in a drug product approved by the federal Food and Drug
Administration; Delta 6 cis or trans tetrahydrocannabinol and their
optical isomers; and Delta 3,4 cis or trans tetrahydrocannabinol
and its optical isomers. Since nomenclature of these substances is
not internationally standardized, compounds of these structures
shall be included regardless of the numerical designation of atomic
positions covered; (16) 3,4-methylenedioxy amphetamine; (17)
5-methoxy-3, 4-methylenedioxy amphetamine; (18) 3,4,5-trimethoxy
amphetamine; (19) N-ethyl-3-piperidyl benzilate; (20)
N-methyl-3-piperidyl benzilate; (21) thiophene analog of
phencyclidine. Trade and other names shall include, but are not
limited to: 1-(1-(2-thienyl)-cyclohexyl)-piperidine;
2-thienylanalog of phencyclidine; TPCP; and TCP; (22)
2,5-dimethoxyamphetamine. Trade and other names shall include, but
are not limited to: 2,5-dimethoxy-a-methylphenethylamine; and
2,5-DMA; (23) hashish or concentrated cannabis having more than
three-tenths of one percent of tetrahydrocannabinol; (24)
Parahexyl. Trade and other names shall include, but are not
limited to: 3-Hexyl-1-hydroxy-7,8,9,10-tetrahydro-6,6,
9-trimethyl-6H-dibenzo(b,d)pyran; and synhexyl; (25) Ethylamine
analog of phencyclidine. Trade and other names shall include, but
are not limited to: N-ethyl-1-phenylcyclohexylamine;
(1-phenylcyclohexyl)ethylamine; N-(1-phenylcyclohexyl)ethylamine; cyclohexamine; and PCE; (26) Pyrrolidine analog of phencyclidine.

Trade and other names shall include, but are not limited to:

1-(1-phenylcyclohexyl)-pyrrolidine; PCPy; and PHP; (27)
3,4-methylenedioxymethamphetamine (MDMA), its optical, positional, and geometric isomers, salts, and salts of isomers; and (28) Phenethylamine. Trade and other names shall include, but are not limited to: 4-bromo-2,5-dimethoxyphenethylamine; 2-CB; Venus; Bromo; Erox; and Nexus.

(d) 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) Mecloqualone; and (2) methaqualone.

(e) 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) Fenethylline; and (2) N-ethylamphetamine.

(f) Gamma hydroxy butyrate (GHB).

(g) Any controlled substance analogue to the extent intended for human consumption.

Schedule II

(a) Any of the following substances except those narcotic
drugs listed in other schedules whether produced directly or
indirectly by extraction from substances of vegetable origin,
independently by means of chemical synthesis, or by combination of
extraction and chemical synthesis:

(1) Opium and opiate, and any salt, compound, derivative,
or preparation of opium or opiate, excluding apomorphine,
buprenorphine, nalbuphine, nalmefene, naloxone, and naltrexone and
their salts, but including the following: (i) Raw opium; (ii) opium
extracts; (iii) opium fluid extracts; (iv) powdered opium; (v)
granulated opium; (vi) tincture of opium; (vii) codeine; (viii)
ethylmorphine; (ix) etorphine hydrochloride; (x) dihydrocodeinone
which is also known as hydrocodone; (xi) hydromorphone; (xii)
metopon; (xiii) morphine; (xiv) oxycodone; (xv) oxymorphone; and
(xvi) thebaine;

(2) Any salt, compound, derivative, or preparation
thereof which is chemically equivalent to or identical with any of
the substances referred to in subdivision (1) of this subdivision,
except that these substances shall not include the isoquinoline
alkaloids of opium;

(3) Opium poppy and poppy straw;

(4) Coca leaves and any salt, compound, derivative, or
preparation of coca leaves, and any salt, compound, derivative, or
preparation thereof which is chemically equivalent to or identical
with any of these substances, including cocaine and its salts,
optical isomers, and salts of optical isomers, except that the
substances shall not include decocainized coca leaves or
extractions which do not contain cocaine or ecgonine; and

(5) Concentrate of poppy straw, the crude extract of
poppy straw in either liquid, solid, or powder form which contains
the phenanthrine alkaloids of the opium poppy.

(b) Unless specifically excepted or unless in another
schedule any of the following opiates, including their isomers,
esters, ethers, salts, and salts of their isomers, esters, and
ethers whenever the existence of such isomers, esters, ethers, and
salts is possible within the specific chemical designation,
dextrorphan and levopropoxyphene excepted: (1) Alphaprodine; (2)
anileridine; (3) bezitramide; (4) diphenoxylate; (5) fentanyl; (6)
levo-methadone; (7) levomethorphan; (8) levorphanol; (9) metazocine;
(10) methadone; (11) methadone-Intermediate,
4-cyano-2-dimethylamino-4, 4-diphenyl butane; (12)
moramide-Intermediate, 2-methyl-3-morpholino-1,
1-diphenyl-propane-carboxylic acid; (13) pethidine or meperidine;
(14) pethidine-Intermediate-A, 4-cyano-1-methyl-4-phenylpiperidine;
(15) pethidine-Intermediate-B,
ethyl-4-phenylpiperidine-4-carboxylate; (16)
pethidine-Intermediate-C, 1-methyl-4-phenylpiperidine-4-carboxylic
acid; (17) phenazocine; (18) piminodine; (19) racemethorphan; (20)
racemorphan; (21) dihydrocodeine; (22) bulk dextropropoxyphene in
nondosage forms; (23) sufentanil; (24) alfentanil; and (25)
levo-alphacetylmethadol which is also known as
levo-alpha-acetylmethadol, levomethadyl acetate, and LAAM.

(c) Any material, compound, mixture, or preparation which
contains any quantity of the following substances having a
potential for abuse associated with a stimulant effect on the
central nervous system: (1) Amphetamine, its salts, optical
isomers, and salts of its optical isomers; (2) phenmetrazine and
its salts; (3) methamphetamine, its salts, isomers, and salts of its isomers; and (4) methylphenidate.

(d) Any material, compound, mixture, or preparation which contains any quantity of the following substances having a potential for abuse associated with 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 designations:

(1) Amobarbital;  (2) secobarbital; (3) pentobarbital; (4) phencyclidine; and (5) glutethimide.

(e) Hallucinogenic substances known as: (1) Nabilone. Another name for nabilone is (+)-trans-3-(1,1-dimethylheptyl)-6,6a,7,8,10,10a-hexahydro-1-hydroxy-6,6-dimethyl-9H-dibenzo(b,d)pyran-9-one.

(f) 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: Phenylacetone. Trade and other names shall include, but are not limited to: Phenyl-2-propanone; P2P; benzyl methyl ketone; and methyl benzyl ketone; or (2) immediate precursors to phencyclidine, PCP: (i) 1-phenylcyclohexylamine; or (ii) 1-piperidinocyclohexanecarbonitrile, PCC.

Schedule III

(a) Any material, compound, mixture, or preparation which contains any quantity of the following substances having a potential for abuse associated with a stimulant effect on the central nervous system, including their salts, isomers, whether

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optical, position, or geometric, and salts of such isomers whenever
the existence of such salts, isomers, and salts of isomers is
possible within the specific chemical designation: (1)
Benzphetamine; (2) chlortermine; (3) chlortermine; and (4)
phendimetrazine.

(b) Any material, compound, mixture, or preparation which
contains any quantity of the following substances having a
potential for abuse associated with a depressant effect on the
central nervous system: (1) Any substance which contains any
quantity of a derivative of barbituric acid or any salt of a
derivative of barbituric acid, except those substances which are
specifically listed in other schedules of this section; (2)
chlorhexadol; (3) lysergic acid; (4) lysergic acid amide; (5)
methyprylon; (6) sulfondiethylmethane; (7) sulfonethylmethane; (8)
sulfonmethane; (9) nalorphine; (10) any compound, mixture, or
preparation containing amobarbital, secobarbital, pentobarbital, or
any salt thereof and one or more other active medicinal ingredients
which are not listed in any schedule; (11) any suppository dosage
form containing amobarbital, secobarbital, pentobarbital, or any
salt of any of these drugs and approved by the Food and Drug
Administration for marketing only as a suppository; and (12)
tiletamine and zolazepam or any salt thereof. Trade or other names
for a tiletamine-zolazepam combination product shall include, but
not be limited to: telazol. Trade or other names for tiletamine
shall include, but not be limited to:
2-(ethylamino)-2-(2-thienyl)-cyclohexanone. Trade or other names
for zolazepam shall include, but not be limited to:
4-(2-fluorophenyl)-6, 8-dihydro-1,3,8-trimethylpyrazolo-(3,4-e)
(1,4)-diazepin-7(1H)-one, and flupyrazapon.

(c) Any material, compound, mixture, or preparation containing limited quantities of any of the following narcotic drugs, or any salts calculated as the free anhydrous base or alkaloid, in limited quantities as set forth below:

(1) Not more than one and eight-tenths grams of codeine per one hundred milliliters or not more than ninety milligrams per dosage unit, with an equal or greater quantity of an isoquinoline alkaloid of opium;

(2) Not more than one and eight-tenths grams of codeine per one hundred milliliters or not more than ninety milligrams per dosage unit, with one or more active, nonnarcotic ingredients in recognized therapeutic amounts;

(3) Not more than three hundred milligrams of dihydrocodeinone which is also known as hydrocodone per one hundred milliliters or not more than fifteen milligrams per dosage unit, with a fourfold or greater quantity of an isoquinoline alkaloid of opium;

(4) Not more than three hundred milligrams of dihydrocodeinone which is also known as hydrocodone per one hundred milliliters or not more than fifteen milligrams per dosage unit, with one or more active, nonnarcotic ingredients in recognized therapeutic amounts;

(5) Not more than one and eight-tenths grams of dihydrocodeine per one hundred milliliters or not more than ninety milligrams per dosage unit, with one or more active, nonnarcotic ingredients in recognized therapeutic amounts;

(6) Not more than three hundred milligrams of
ethylmorphine per one hundred milliliters or not more than fifteen
milligrams per dosage unit, with one or more active, nonnarcotic
ingredients in recognized therapeutic amounts;

(7) Not more than five hundred milligrams of opium per
one hundred milliliters or per one hundred grams, or not more than
twenty-five milligrams per dosage unit, with one or more active,
nonnarcotic ingredients in recognized therapeutic amounts; and

(8) Not more than fifty milligrams of morphine per one
hundred milliliters or per one hundred grams with one or more
active, nonnarcotic ingredients in recognized therapeutic amounts.

(d) Any anabolic steroid, which shall include any
material, compound, mixture, or preparation containing any quantity
of the following substances, including its salts, isomers, and
salts of isomers whenever the existence of such salts of isomers is
possible within the specific chemical designation: (1) Boldenone;
(2) chlorotestosterone (4-chlortestosterone); (3) clostebol; (4)
dehydrochlormethyltestosterone; (5) dihydrotestosterone
(4-dihydrotestosterone); (6) drostanolone; (7) ethylestrenol; (8)
fluoxymesterone; (9) formebulone (formebolone); (10) mesterolone;
(11) methandienone; (12) methandranone; (13) methandriol; (14)
methandrostenolone; (15) methenolone; (16) methyltestosterone; (17)
mibolerone; (18) nandrolone; (19) norethandrolone; (20)
oxandrolone; (21) oxymesterone; (22) oxymetholone; (23) stanolone;
(24) stanozolol; (25) testolactone; (26) testosterone; (27)
trenbolone; and (28) any salt, ester, or isomer of a drug or
substance described or listed in this subdivision if the salt,
ester, or isomer promotes muscle growth.

(e) Hallucinogenic substances known as: (1) Dronabinol,
synthetic, in sesame oil and encapsulated in a soft gelatin capsule
in a Food and Drug Administration approved drug product. Some
other names for dronabinol are
(6aR-trans)-6a,7,8,10a-tetrahydro-6,6,9-trimethyl-
3-pentyl-6H-dibenzo(b,d)pyran-1-ol
or
(-)-delta-9-(trans)-tetrahydrocannabinol.

(f) Ketamine, its salts, isomers, and salts of isomers.

Some other names for ketamine:
(+)-2-(2-chlorophenyl)-2-(methylamino)-cyclohexanone.

Schedule IV

(a) Any material, compound, mixture, or preparation which
contains any quantity of the following substances, 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) Barbital; (2) chloral betaine;
(3) chloral hydrate; (4) chlor Diazepoxide, but not including librax
(chlordiazepoxide hydrochloride and clindinium bromide) or menrium
(chlordiazepoxide and water soluble esterified estrogens); (5)
clonazepam; (6) clorazepate; (7) diazepam; (8) ethchlorvynol; (9)
ethinamate; (10) flurazepam; (11) mebutamate; (12) meprobamate;
(13) methohexital; (14) methylphenobarbital; (15) oxazepam; (16)
paraldehyde; (17) petrichloral; (18) phenobarbital; (19) prazepam;
(20) alprazolam; (21) bromazepam; (22) camazepam; (23) clonazepam;
(24) clotiazepam; (25) cloxazolam; (26) delorazepam; (27)
estazolam; (28) ethyl loflazepate; (29) fludiazepam; (30)
flunitrazepam; (31) halazepam; (32) haloxazolam; (33) ketazolam;
(34) lorazepam; (35) lormetazepam; (36) nimezapam; (37) medazepam;
(38) nitrazepam; (39) nordiazepam; (40) oxazolam;
(42) pinazepam; (43) temazepam; (44) tetrazepam; (45) triazolam;
(46) midazolam; (47) quazepam; and (48) zolpidem.

(b) Any material, compound, mixture, or preparation which
contains any quantity of the following substance, including its
salts, isomers, whether optical, position, or geometric, and salts
of such isomers, whenever the existence of such salts, isomers, and
salts of isomers is possible: Fenfluramine.

(c) 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 their
salts, isomers, whether optical, position, or geometric, and salts
of such isomers whenever the existence of such salts, isomers, and
salts of isomers is possible within the specific chemical
designation: (1) Diethylpropion; (2) phentermine; (3) pemoline,
including organometallic complexes and chelates thereof; (4)
mazindol; (5) pipradrol; (6) SPA,((-)-1-dimethylamino-1,2-diphenylethane); (7) cathine. Another
name for cathine is ((+)-norpseudoephedrine); (8) fencamfamin; (9)
fenproporex; and (10) mefenorex.

(d) Unless specifically excepted or unless listed in
another schedule, any material, compound, mixture, or preparation
which contains any quantity of the following narcotic drugs, or
their salts or isomers calculated as the free anhydrous base or
alkaloid, in limited quantities as set forth below: (1)
Propoxyphene; and (2) not more than one milligram of difenoxin and
not less than twenty-five micrograms of atropine sulfate per dosage
unit.
(e) Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substance, including its salts: Pentazocine.

(f) Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substance, including its salts, isomers, and salts of such isomers: Butorphanol.

(g)(1) Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation which contains any quantity of the following substance, including its salts, optical isomers, and salts of such optical isomers: Ephedrine.

(2) The following drug products containing ephedrine, its salts, optical isomers, and salts of such optical isomers are excepted from subdivision (g)(1) of Schedule IV if they may lawfully be sold over the counter without a prescription under the Federal Food, Drug, and Cosmetic Act; are labeled and marketed in a manner consistent with the pertinent OTC Tentative Final or Final Monograph; are manufactured and distributed for legitimate medicinal use in a manner that reduces or eliminates the likelihood of abuse; and are not marketed, advertised, or labeled for the indication of stimulation, mental alertness, weight loss, muscle enhancement, appetite control, or energy:

(A) Solid oral dosage forms, including soft gelatin capsules, that combine active ingredients in the following ranges for each dosage unit:

(i) Not less than one hundred milligrams nor more than
one hundred thirty milligrams of theophylline and not less than
twelve and five-tenths milligrams nor more than twenty-four
milligrams of ephedrine;

(ii) Not less than sixty milligrams nor more than one
hundred milligrams of theophylline, not less than twelve and
five-tenths milligrams nor more than twenty-four milligrams of
ephedrine, and not less than two hundred milligrams nor more than
four hundred milligrams of guaifenesin;

(iii) Not less than twelve and five-tenths milligrams nor
more than twenty-five milligrams of ephedrine and not less than two
hundred milligrams nor more than four hundred milligrams of
guaifenesin; and

(iv) Not more than eight milligrams of phenobarbital in
combination with the ingredients of subdivision (g)(2)(A)(i) or
(g)(2)(A)(ii) of Schedule IV;

(B) Liquid oral dosage forms that combine active
ingredients in the following ranges for each five-milliliter dose:

(i) Not more than forty-five milligrams of theophylline,
not more than thirty-six milligrams of ephedrine, not more than one
hundred milligrams of guaifenesin, and not more than twelve
milligrams of phenobarbital; and

(ii) Not more than five milligrams of phenylephrine, not
more than five milligrams of ephedrine, not more than two
milligrams of chlorpheniramine, not more than ten milligrams of
dextromethorphan, not more than forty milligrams of ammonium
chloride, and not more than five one-thousandths of a milligram of
ipecac fluid extract; and

(C) Anorectal preparations containing less than five
Schedule V

(a) Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation containing any of the following narcotic drug and its salts: (1) Buprenorphine.

(b) Any compound, mixture, or preparation containing any of the following limited quantities of narcotic drugs or salts calculated as the free anhydrous base or alkaloid, which shall include 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 two hundred milligrams of codeine per one hundred milliliters or per one hundred grams;

(2) Not more than one hundred milligrams of dihydrocodeine per one hundred milliliters or per one hundred grams;

(3) Not more than one hundred milligrams of ethylmorphine per one hundred milliliters or per one hundred grams;

(4) Not more than two and five-tenths milligrams of diphenoxylate and not less than twenty-five micrograms of atropine sulfate per dosage unit;

(5) Not more than one hundred milligrams of opium per one hundred milliliters or per one hundred grams; and

(6) Not more than five-tenths milligram of difenoxin and not less than twenty-five micrograms of atropine sulfate per dosage unit.
Sec. 10. Section 81-2,147.06, Revised Statutes Supplement, 2000, is amended to read:

81-2,147.06. (1) The duty of enforcing the Nebraska Seed Law and carrying out such law and requirements shall be vested in
the director. It shall be the duty of the director:

(a) To sample, inspect, make analysis of, and test agricultural, vegetable, and flower seed sold within this state for sowing purposes at such time and place and to such extent as he or she may deem necessary to determine whether such agricultural, vegetable, or flower seed is in compliance with the Nebraska Seed Law and to notify promptly the persons who sold the seed of any violation;

(b) To adopt and promulgate rules and regulations in compliance with the Administrative Procedure Act as are specifically authorized in the Nebraska Seed Law governing the method of sampling, inspecting, analyzing, testing, and examining agricultural, vegetable, and flower seed and the tolerances to be followed in the administration of the law, which shall be in general accord with officially prescribed practice in interstate commerce, and such other rules and regulations as may be necessary to secure the efficient enforcement and full intent of such law;

(c) To adopt and promulgate rules and regulations in compliance with the Administrative Procedure Act adding to or subtracting from the primary noxious weed seeds list, the prohibited noxious weed seeds list, and the restricted noxious weed seeds list, as defined in section 81-2,147.01, whenever the director finds that a noxious weed seed should or should not be within one of these lists.
having no more than three-tenths of one percent tetrahydrocannabinol shall not be designated as a noxious weed seed;

(d) To adopt and promulgate rules and regulations in compliance with the Administrative Procedure Act establishing reasonable standards of germination for agricultural, vegetable, and flower seed; and

(e) To adopt and promulgate rules and regulations in compliance with the Administrative Procedure Act to establish, add to, or subtract from the seeds listed in subdivision (2)(i) of section 81-2,147.02 and for which the tetrazolium (TZ) test may be employed as the official test to indicate the potential viability of the seed.

(2) For the purpose of carrying out the law, the director may:

(a) Enter upon any public or private premises during regular business hours in order to have access to seeds and the records connected with such seeds subject to the law and the rules and regulations adopted and promulgated under such law and enter any truck or other conveyor by land, water, or air at any time when the conveyor is accessible for the same purpose;

(b) Issue and enforce a written or printed stop-sale order to the owner or custodian of any lot of agricultural, vegetable, or flower seed which the director finds is in violation of any of the provisions of the law or rules and regulations adopted and promulgated under such law, which order shall prohibit further sale, conditioning, and movement of such seed, except on approval of the enforcing officer, until such officer has evidence
that the law has been complied with and he or she has issued a release from the stop-sale order of such seed. With respect to seed which has been denied sale, conditioning, or movement as provided in this subdivision, the owner or custodian of such seed shall have the right to appeal from such order in accordance with the Administrative Procedure Act, praying for a judgment as to the justification of such order and for the discharge of such seed from the order prohibiting the sale, conditioning, or movement in accordance with the findings of the court. This subdivision shall not be construed as limiting the right of the director to proceed as authorized by other sections of the law;

c) Establish and maintain or make provision for seed-testing facilities, employ qualified persons, and incur such expenses as may be necessary to comply with the law or rules and regulations adopted and promulgated under the law;

d) Make or provide for making purity, weed seed, tetrazolium (TZ), germination, and other tests of seed as established in rules and regulations and recommended by rule of the Association of Official Seed Analysts for persons on request, adopt and promulgate rules and regulations in compliance with the Administrative Procedure Act governing such testing, and fix and collect charges for the tests made, which charges shall not exceed the cost of such tests. All fees shall be remitted to the state treasury and by the State Treasurer placed in the Nebraska Seed Administrative Cash Fund;

e) Cooperate with the United States Department of Agriculture and other agencies in seed law enforcement; and

(f) Cooperate and enter into agreements with any person
necessary to carry out the purpose of the law.

Sec. 11. Original sections 2-954, 16-230, and 17-563, Reissue Revised Statutes of Nebraska, and sections 28-401, 28-405, and 81-2,147.06, Revised Statutes Supplement, 2000, are repealed.