

## 103RD GENERAL ASSEMBLY State of Illinois 2023 and 2024 HB2306

Introduced 2/14/2023, by Rep. Lakesia Collins

## SYNOPSIS AS INTRODUCED:

See Index

Amends the Physician Assistant Practice Act of 1987. Changes the definition of "physician assistant", "physician assistant practice", "board", and "collaborating physician". Removes the definition of "disciplinary board" and changes references from the "disciplinary board" to the Illinois State Medical Board throughout the Act. Provides that a physician assistant shall be deemed by law to possess the ability to prescribe, dispense, order, administer, and procure drugs and medical devices without delegation of such authority by a physician. Provides that such ability shall include prescribing of Schedule II, III, IV, and V controlled substances. Provides that to prescribe Schedule II, III, IV, or V controlled substances under the Act, a physician assistant shall obtain a mid-level practitioner controlled substances licenses. Provides that when a written collaboration agreement is required under the Act, delegation of prescriptive authority by a physician is not required. Provides that a physician assistant who files with the Department of Financial and Professional Regulation a notarized attestation of completion of at least 250 hours of continuing education or training and at least 2,000 hours of clinical experience after first attaining national certification shall not require a written collaborative agreement. Provides the specified scope of practice of a physician assistant with optimal practice authority. Provides that a physician assistant shall be able to hold more than one professional position. Makes changes in provisions concerning the physician assistant title, collaboration requirements, and the written collaborative agreement. Makes other changes and corresponding changes to the Act and to the Illinois Controlled Substances Act.

LRB103 05245 AMQ 50263 b

1 AN ACT concerning regulation.

## Be it enacted by the People of the State of Illinois, represented in the General Assembly:

- 4 Section 5. The Physician Assistant Practice Act of 1987 is
- 5 amended by changing Sections 4, 5.5, 6, 7, 7.5, 7.7, 17, 21,
- 6 22.2, 22.3, 22.5, 22.6, 22.7, 22.8, 22.9, and 22.10 and by
- 7 adding Sections 7.8 and 7.9 as follows:
- 8 (225 ILCS 95/4) (from Ch. 111, par. 4604)
- 9 (Section scheduled to be repealed on January 1, 2028)
- 10 Sec. 4. Definitions. In this Act:
- 11 1. "Department" means the Department of Financial and
- 12 Professional Regulation.
- 13 2. "Secretary" means the Secretary of Financial and
- 14 Professional Regulation.
- 3. "Physician assistant" means any person not holding an
- 16 active license or permit issued by the Department pursuant to
- 17 the Medical Practice Act of 1987 who has been certified as a
- 18 physician assistant by the National Commission on the
- 19 Certification of Physician Assistants or equivalent successor
- 20 agency. and performs procedures in collaboration with a
- 21 physician as defined in this Act. A physician assistant may
- 22 perform such procedures within the specialty of the
- 23 collaborating physician, except that such physician shall

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exercise such direction, collaboration, and control over such physician assistants as will assure that patients shall receive quality medical care. Physician assistants shall be capable of performing a variety of tasks within the specialty of medical care in collaboration with a physician. Collaboration with the physician assistant shall not be construed to necessarily require the personal presence of the collaborating physician at all times at the place where services are rendered, as long as there is communication available for consultation by radio, telephone or telecommunications within established guidelines as determined by the physician/physician assistant team. The collaborating physician may delegate tasks and duties to the physician assistant. Delegated tasks or duties shall be consistent with physician assistant education, training, and experience. The delegated tasks or duties shall be specific to the practice setting and shall be implemented and reviewed under a written collaborative agreement established by the physician physician/physician assistant team. A physician assistant, acting as an agent of the physician, shall be permitted to transmit the collaborating physician's orders as determined by the institution's by-laws, policies, procedures, or job description within which the physician/physician assistant team practices. Physician assistants shall practice only in accordance with a written collaborative agreement.

Any person who holds an active license or permit issued

1	pursuant to the Medical Practice Act of 1987 shall have that
2	license automatically placed into inactive status upon
3	issuance of a physician assistant license. Any person who
4	holds an active license as a physician assistant who is issued
5	a license or permit pursuant to the Medical Practice Act of
6	1987 shall have his or her physician assistant license
7	automatically placed into inactive status.
8	3.5. "Physician assistant practice" means the performance
9	of any legal medical service for which the physician assistant
10	has been prepared by the physician assistant's education,
11	training, and experience and is competent to perform as
12	determined by the practice through employment agreement or
13	credentialing and privileging systems of licensed facilities.
14	Medical and surgical services provided by physician assistants
15	<pre>include, but are not limited to:</pre>
16	(A) obtaining and performing comprehensive health
17	histories and physical examinations;
18	(B) evaluating, diagnosing, managing, and providing
19	<pre>medical treatment;</pre>
20	(C) ordering, performing, and interpreting diagnostic
21	studies and therapeutic procedures;
22	(D) educating patients on health promotion and disease
23	<pre>prevention;</pre>
24	(E) providing consultation upon request;
25	<pre>(F) writing medical orders;</pre>
26	(G) prescribing, dispensing, ordering, administering,

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(H) assisting in surgery. procedures within the specialty of the collaborating physician. Physician assistants shall be capable of performing a variety of tasks within the specialty of medical care of the collaborating physician. Collaboration with the physician assistant shall not be construed to necessarily require the personal presence of the collaborating physician at all times at the place where services are rendered, as long as there is communication available for consultation by radio, telephone, telecommunications, or electronic communications. The collaborating physician may delegate tasks and duties to the physician assistant. Delegated tasks or duties shall be consistent with physician assistant education, training, and experience. The delegated tasks or duties shall be specific to the practice setting and shall be implemented and reviewed under a written collaborative agreement established by the physician or physician/physician assistant team. A physician assistant shall be permitted to transmit the collaborating physician's orders as determined by the institution's bylaws, policies, or procedures or the job description within which the physician/physician assistant team practices. Physician assistants shall practice only in accordance with a written collaborative agreement, except as provided in Section 7.5 of this Act.

- 1 4. "Board" means the <u>Illinois State Medical Board</u> <u>Medical</u>
- 2 Licensing Board constituted under the Medical Practice Act of
- 3 <del>1987</del>.
- 4 5. (Blank). "Disciplinary Board" means the Medical
- 5 Disciplinary Board constituted under the Medical Practice Act
- 6 <del>of 1987.</del>
- 7 6. "Physician" means a person licensed to practice
- 8 medicine in all of its branches under the Medical Practice Act
- 9 of 1987.
- 7. "Collaborating physician" means the physician who,
- 11 within his or her specialty and expertise, may delegate a
- 12 variety of tasks and procedures to the physician assistant.
- 13 Such tasks and procedures shall be delegated in accordance
- 14 with a written collaborative agreement when such agreement is
- 15 required under this Act.
- 16 8. (Blank).
- 9. "Address of record" means the designated address
- 18 recorded by the Department in the applicant's or licensee's
- 19 application file or license file maintained by the
- 20 Department's licensure maintenance unit.
- 21 10. "Hospital affiliate" means a corporation, partnership,
- 22 joint venture, limited liability company, or similar
- organization, other than a hospital, that is devoted primarily
- 24 to the provision, management, or support of health care
- 25 services and that directly or indirectly controls, is
- 26 controlled by, or is under common control of the hospital. For

- 1 the purposes of this definition, "control" means having at
- least an equal or a majority ownership or membership interest.
- 3 A hospital affiliate shall be 100% owned or controlled by any
- 4 combination of hospitals, their parent corporations, or
- 5 physicians licensed to practice medicine in all its branches
- 6 in Illinois. "Hospital affiliate" does not include a health
- 7 maintenance organization regulated under the Health
- 8 Maintenance Organization Act.
- 9 11. "Email address of record" means the designated email
- 10 address recorded by the Department in the applicant's
- 11 application file or the licensee's license file, as maintained
- by the Department's licensure maintenance unit.
- 13 (Source: P.A. 99-330, eff. 1-1-16; 100-453, eff. 8-25-17.)
- 14 (225 ILCS 95/5.5)
- 15 (Section scheduled to be repealed on January 1, 2028)
- 16 Sec. 5.5. Billing. A physician assistant may shall not be
- 17 allowed to personally bill patients and or in any way charge
- 18 for services. The employer of a physician assistant may bill
- 19 and charge for services rendered by the physician assistant.
- 20 All claims for services rendered by the physician assistant
- 21 shall be submitted using the physician assistant's national
- 22 provider identification number as the rendering provider, with
- 23 the exception of when optional billing provisions, such as
- 24 incident to, split, or shared visit billing, are being used
- 25 whenever appropriate. Payment for services rendered by a

- 1 physician assistant shall be made to his or her employer if the
- 2 payor would have made payment had the services been provided
- 3 by a physician licensed to provide medicine in all of its
- 4 branches.
- 5 (Source: P.A. 100-453, eff. 8-25-17; 100-559, eff. 12-8-17.)
- 6 (225 ILCS 95/6) (from Ch. 111, par. 4606)
- 7 (Section scheduled to be repealed on January 1, 2028)
- 8 Sec. 6. Physician assistant title.
- 9 (a) No physician assistant shall use the title of doctor
- 10 or, physician, or associate with his or her name or any other
- 11 term that would indicate to other persons that he or she is
- 12 qualified to engage in the general practice of medicine.
- 13 (b) A physician assistant shall verbally identify himself
- or herself as a physician assistant, including specialty
- 15 certification, when applicable, to each patient.
- 16 (c) Nothing in this Act shall be construed to relieve a
- 17 physician assistant of the professional or legal
- 18 responsibility for the care and treatment of persons attended
- 19 by him or her.
- 20 (d) (Blank). The collaborating physician shall file with
- 21 the Department notice of employment, discharge, or
- 22 collaboration with a physician assistant within 60 days of
- 23 employment, discharge, or assumption of collaboration with a
- 24 physician assistant. Nothing in this Section shall prevent a
- 25 physician assistant from beginning his or her employment

- 1 before the notice of employment or collaboration has been
- 2 filed.
- 3 (Source: P.A. 102-735, eff. 1-1-23.)
- 4 (225 ILCS 95/7) (from Ch. 111, par. 4607)
- 5 (Section scheduled to be repealed on January 1, 2028)
- 6 Sec. 7. Collaboration requirements.
- 7 (a) A written collaborative agreement is required for all
- 8 physician assistants engaged in clinical practice prior to
- 9 <u>meeting the requirements of Section 7.9, except for physician</u>
- 10 assistants who practice in a hospital, hospital affiliate, or
- 11 ambulatory surgical treatment center as provided in Section
- 12 7.7.
- 13 (b) A collaborating physician shall determine the number
- of physician assistants to collaborate with, provided the
- 15 physician is able to provide adequate collaboration as
- outlined in the written collaborative agreement required under
- 17 Section 7.5 of this Act and consideration is given to the
- nature of the physician's practice, complexity of the patient
- 19 population, and the experience of each physician assistant. A
- 20 collaborating physician may collaborate with a maximum of 7
- 21 <u>full-time equivalent physician assistants as described in</u>
- 22 Section 54.5 of the Medical Practice Act of 1987. As used in
- 23 this Section, "full-time equivalent" means the equivalent of
- 24 40 hours per week per individual. Physicians and physician
- 25 assistants who work in a hospital, hospital affiliate, or

- ambulatory surgical treatment center as defined by Section 7.7 of this Act are exempt from the collaborative ratio restriction requirements of this Section. A physician assistant shall be able to hold more than one professional position. A collaborating physician shall file a notice of collaboration of each physician assistant according to the rules of the Department.
  - (c) A physician assistant shall be able to hold more than one professional position.
  - (d) Physician assistants shall collaborate only with physicians as defined in this Act who are engaged in clinical practice, or in clinical practice in public health or other community health facilities.
  - (e) Nothing in this Act shall be construed to limit the delegation of tasks or duties by a physician to a nurse or other appropriately trained personnel.
  - (f) Nothing in this Act shall be construed to prohibit the employment of physician assistants by a hospital, nursing home or other health care facility where such physician assistants function with under a collaborating physician.
  - (g) A physician assistant may be employed by a practice group or other entity employing multiple physicians at one or more locations. In that case, one of the physicians practicing at a location shall be designated the collaborating physician. The other physicians with that practice group or other entity who practice in the same general type of practice or specialty

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- as the collaborating physician may collaborate with the physician assistant with respect to their patients.
- (h) (b) A physician assistant licensed in this State, or licensed or authorized to practice in any other U.S. jurisdiction or credentialed by his or her federal employer as a physician assistant, who is responding to a need for medical care created by an emergency or by a state or local disaster may render such care that the physician assistant is able to provide without collaboration as it is defined in this Section
- 11 <u>(i)</u> Any physician who collaborates with a physician assistant providing medical care in response to such an emergency or state or local disaster shall not be required to meet the requirements set forth in this Section for a collaborating physician.
- 16 (Source: P.A. 100-453, eff. 8-25-17; 100-605, eff. 1-1-19.)

or with such collaboration as is available.

- 17 (225 ILCS 95/7.5)
- 18 (Section scheduled to be repealed on January 1, 2028)
- 19 Sec. 7.5. Written collaborative agreements; prescriptive authority.
  - (a) A written collaborative agreement is required for all physician assistants to practice in the State, except as provided in Section 7.7 and Section 7.9 of this Act. When a written collaborative agreement is required under this Act,
- 25 the following shall apply:

(1) A written collaborative agreement shall describe 1 the working relationship of the physician assistant with 2 3 collaborating physician and shall describe categories of care, treatment, or procedures to 4 5 provided by the physician assistant. The written 6 collaborative agreement shall promote the exercise of 7 judgment by the physician <del>professional</del> 8 commensurate with his or her education and experience. The 9 services to be provided by the physician assistant shall 10 be services that the collaborating physician is authorized 11 to and generally provides to his or her patients in the 12 normal course of his or her clinical medical practice. The written collaborative agreement need not describe 13 14 exact steps that a physician assistant must take with 15 respect to each specific condition, disease, or symptom 16 but must specify which authorized procedures require the 17 presence of the collaborating physician as the procedures are being performed. The relationship under a written 18 19 collaborative agreement shall not be construed to require the personal presence of a physician at the place where 20 services are rendered. Methods of communication shall be 21 22 available for consultation with the collaborating 23 physician in person or by telecommunications or electronic communications as set forth in the written collaborative 24 agreement. For the purposes of this Act, "generally 25 26 provides to his or her patients in the normal course of his

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Τ	or ner clinical medical practice" means services, not
2	specific tasks or duties, the collaborating physician
3	routinely provides individually or through delegation to
4	other persons so that the physician has the experience and
5	ability to collaborate and provide consultation.
6	(2) (Blank). The written collaborative agreement shall
7	be adequate if a physician does each of the following:
8	(A) Participates in the joint formulation and
9	joint approval of orders or guidelines with the
10	physician assistant and he or she periodically reviews
11	such orders and the services provided patients under
12	such orders in accordance with accepted standards of
13	medical practice and physician assistant practice.
14	(B) Provides consultation at least once a month.
15	(3) A copy of the signed, written collaborative
16	agreement must be available to the Department upon request
17	from both the physician assistant and the collaborating
18	<del>physician</del> .
19	(4) A physician assistant shall inform each
20	collaborating physician of all written collaborative
21	agreements he or she has signed and provide a copy of these
22	to any collaborating physician upon request.
23	(b) To prescribe Schedule II, III, IV, or V controlled
24	substances under this Section, a physician assistant must

obtain a mid-level practitioner controlled substances license.

A collaborating physician may, but is not required to,

delegate prescriptive authority to a physician assistant as part of a written collaborative agreement. This authority may, but is not required to, include prescription of, selection of, orders for, administration of, storage of, acceptance of samples of, and dispensing medical devices, over the counter medications, legend drugs, medical gases, and controlled substances categorized as Schedule II through V controlled substances, as defined in Article II of the Illinois Controlled Substances Act, and other preparations, including, but not limited to, botanical and herbal remedies. The collaborating physician must have a valid, current Illinois controlled substance license and federal registration with the Drug Enforcement Administration to delegate the authority to prescribe controlled substances.

(1) To prescribe Schedule II, III, IV, or V controlled substances under this Section, a physician assistant must obtain a mid level practitioner controlled substances license. Medication orders issued by a physician assistant shall be reviewed periodically by the collaborating physician.

(2) The collaborating physician shall file with the Department notice of delegation of prescriptive authority to a physician assistant and termination of delegation, specifying the authority delegated or terminated. Upon receipt of this notice delegating authority to prescribe controlled substances, the physician assistant shall be

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eligible to register for a mid-level practitioner controlled substances license under Section 303.05 of the Illinois Controlled Substances Act. Nothing in this Act shall be construed to limit the delegation of tasks or duties by the collaborating physician to a nurse or other appropriately trained persons in accordance with Section 54.2 of the Medical Practice Act of 1987.

(3) In addition to the requirements of this subsection (b), a collaborating physician may, but is not required to, delegate authority to a physician assistant to prescribe Schedule II controlled substances, if all of the following conditions apply:

(A) Specific Schedule II controlled substances by oral dosage or topical or transdermal application may be delegated, provided that the delegated Schedule II controlled substances are routinely prescribed by the collaborating physician. This delegation must identify the specific Schedule II controlled substances by either brand name or generic name. Schedule II controlled substances to be delivered by injection or other route of administration may not be delegated.

(B) (Blank).

(C) Any prescription must be limited to no than a 30-day supply, with any continuation authorized only after prior approval of the collaborating physician.

1	(D) The physician assistant must	<del>-discuss the</del>
2	condition of any patients for whom	a controlled
3	substance is prescribed monthly with the	collaborating
4	<del>physician.</del>	

(E) The physician assistant meets the education requirements of Section 303.05 of the Illinois Controlled Substances Act.

- (c) Nothing in this Act shall be construed to limit the delegation of tasks or duties by a physician to a licensed practical nurse, a registered professional nurse, or other persons. Nothing in this Act shall be construed to limit the method of delegation that may be authorized by any means, including, but not limited to, oral, written, electronic, standing orders, protocols, guidelines, or verbal orders. Nothing in this Act shall be construed to authorize a physician assistant to provide health care services required by law or rule to be performed by a physician. Nothing in this Act shall be construed to authorize the delegation or performance of operative surgery. Nothing in this Section shall be construed to preclude a physician assistant from assisting in surgery.
- (c-5) Nothing in this Section shall be construed to apply to any medication authority, including Schedule II controlled substances of a licensed physician assistant for care provided in a hospital, hospital affiliate, or ambulatory surgical treatment center pursuant to Section 7.7 of this Act or to a

- 1 physician assistant meeting the requirements of Section 7.9 of
- 2 this Act.
- 3 (d) (Blank).
- 4 (e) Nothing in this Section shall be construed to prohibit
- 5 generic substitution.
- 6 (f) Delegation of prescriptive authority by a physician is
- 7 <u>not required under this Section.</u>
- 8 (Source: P.A. 101-13, eff. 6-12-19; 102-558, eff. 8-20-21.)
- 9 (225 ILCS 95/7.7)
- 10 (Section scheduled to be repealed on January 1, 2028)
- 11 Sec. 7.7. Physician assistants in hospitals, hospital
- 12 affiliates, or ambulatory surgical treatment centers.
- 13 (a) A physician assistant may provide services in a
- 14 hospital as defined in the Hospital Licensing Act, a hospital
- 15 affiliate as defined in the University of Illinois Hospital
- 16 Act, or a licensed ambulatory surgical treatment center as
- 17 defined in the Ambulatory Surgical Treatment Center Act
- 18 without a written collaborative agreement pursuant to Section
- 19 7.5 of this Act. A physician assistant must possess clinical
- 20 privileges recommended by the hospital medical staff and
- 21 granted by the hospital or the consulting medical staff
- 22 committee and ambulatory surgical treatment center in order to
- 23 provide services. The medical staff or consulting medical
- 24 staff committee shall periodically review the services of
- 25 physician assistants granted clinical privileges, including

any care provided in a hospital affiliate. A physician assistant practicing under this Section shall have the authority to prescribe, select, order, and administer medications, including controlled substances. Authority may also be granted when recommended by the hospital medical staff and granted by the hospital or recommended by the consulting medical staff committee and ambulatory surgical treatment center to individual physician assistants to select, order, and administer medications, including controlled substances, to provide delineated care. In a hospital, hospital affiliate, or ambulatory surgical treatment center, the attending physician shall determine a physician assistant's role in providing care for his or her patients, except as otherwise provided in the medical staff bylaws or consulting committee policies.

affiliate shall have the authority may be, but are not required to be, granted authority to prescribe Schedule II through V controlled substances when such authority is recommended by the appropriate physician committee of the hospital affiliate and granted by the hospital affiliate. This authority includes may, but is not required to, include prescription of, selection of, orders for, administration of, storage of, acceptance of samples of, and dispensing over-the-counter medications, legend drugs, medical gases, and controlled substances categorized as Schedule II through V

controlled substances, as defined in Article II of the Illinois Controlled Substances Act, and other preparations, including, but not limited to, botanical and herbal remedies.

To prescribe controlled substances under this subsection (a-5), a physician assistant must obtain a mid-level practitioner controlled substance license. Medication orders shall be reviewed periodically by the appropriate hospital affiliate physicians committee or its physician designee.

The hospital affiliate shall file with the Department notice of a grant of prescriptive authority consistent with this subsection (a-5) and termination of such a grant of authority in accordance with rules of the Department. Upon receipt of this notice of grant of authority to prescribe any Schedule II through V controlled substances, the licensed physician assistant may register for a mid-level practitioner controlled substance license under Section 303.05 of the Illinois Controlled Substances Act.

In addition, a hospital affiliate may, but is not required to, grant authority to a physician assistant to prescribe any Schedule II controlled substances if all of the following conditions apply:

(1) specific Schedule II controlled substances by oral dosage or topical or transdermal application may be designated, provided that the designated Schedule II controlled substances are routinely prescribed by physician assistants in their area of certification; this

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<del>other r</del>	oute	of adm	<del>inistr</del>	ation	may no	ot be c	<del>rante</del>	ed;		

- (2) any grant of authority must be controlled substances limited to the practice of the physician assistant;
- (3) any prescription must be limited to no more than a 30 day supply;
- (4) the physician assistant must discuss the condition of any patients for whom a controlled substance is prescribed monthly with the appropriate physician committee of the hospital affiliate or its physician designee; and
- (5) the physician assistant must meet the education requirements of Section 303.05 of the Illinois Controlled Substances Act.
- (b) A physician assistant granted authority to order medications including controlled substances may complete discharge prescriptions provided the prescription is in the name of the physician assistant and the attending or discharging physician.
- (c) Physician assistants practicing in a hospital, hospital affiliate, or an ambulatory surgical treatment center are not required to obtain a mid-level controlled substance

- 1 license to order controlled substances under Section 303.05 of
- the Illinois Controlled Substances Act.
- 3 (d) Delegation of prescriptive authority by a physician is
- 4 not required under this Section.
- 5 (Source: P.A. 100-453, eff. 8-25-17.)
- 6 (225 ILCS 95/7.8 new)
- 7 Sec. 7.8. Prescriptive authority. A physician assistant
- 8 shall be deemed by law to possess the ability to prescribe,
- 9 dispense, order, administer, and procure drugs and medical
- 10 devices without delegation of such authority by a physician.
- 11 Such ability shall include prescribing Schedule II, III, IV,
- and V controlled substances. To prescribe Schedule II, III,
- 13 IV, or V controlled substances under this Act, a physician
- 14 assistant shall obtain a mid-level practitioner controlled
- 15 substances license. When a written collaborative agreement is
- 16 required under this Act, delegation of prescriptive authority
- by a physician is not required.
- 18 (225 ILCS 95/7.9 new)
- 19 Sec. 7.9. Optimal practice authority.
- 20 (a) A physician assistant shall be deemed by law to
- 21 possess the ability to practice without a written
- 22 collaborative agreement as set forth in this Section.
- 23 (b) A physician assistant who files with the Department a
- 24 notarized attestation of completion of at least 250 hours of

1	continuing education or training and at least 2,000 hours of
2	clinical experience after first attaining national
3	certification shall not require a written collaborative
4	agreement. Documentation of successful completion shall be
5	provided to the Department upon request.
6	(c) The scope of practice of a physician assistant with
7	<pre>optimal practice authority includes:</pre>
8	(1) all matters included in subsection (3.5) of
9	Section 4 of this Act;
10	(2) practicing without a written collaborative
11	agreement in all practice settings consistent with this
12	Act;
13	(3) authority to prescribe both legend drugs and
14	Schedule II through V controlled substances; this
15	authority includes prescription of, selection of, orders
16	for, administration of, storage of, acceptance of samples
17	of, and dispensing over-the-counter medications, legend
18	drugs, and controlled substances categorized as any
19	Schedule II through V controlled substances, as defined in
20	Article II of the Illinois Controlled Substances Act, and
21	other preparations, including, but not limited to,
22	botanical and herbal remedies; and
23	(4) authority to obtain a controlled substances
24	license in the State and a federal Drug Enforcement
25	Administration number.

The scope of practice of a physician assistant does not

- 1 <u>include operative surgery. Nothing in this Section shall be</u>
- 2 construed to preclude a physician assistant from assisting in
- 3 surgery or performing other procedures as privileged by the
- 4 physician assistant's employer.
- 5 (d) The Department may adopt rules necessary to administer
- 6 this Section, including, but not limited to, requiring the
- 7 completion of forms and the payment of fees.
- 8 (e) Nothing in this Act shall be construed to authorize a
- 9 physician assistant with optimal practice authority to provide
- 10 <u>health care services required by law or rule to be performed by</u>
- 11 a physician.
- 12 (225 ILCS 95/17) (from Ch. 111, par. 4617)
- 13 (Section scheduled to be repealed on January 1, 2028)
- 14 Sec. 17. Inactive status. Any physician assistant who
- notified the Department in writing on forms prescribed by the
- 16 Department, may elect to place his or her license on an
- inactive status and shall, subject to rules of the Department,
- 18 be excused from payment of renewal fees until he or she
- 19 notifies the Department in writing of his or her intention to
- 20 restore the license. Any person who holds an active license or
- 21 permit issued pursuant to the Medical Practice Act of 1987
- 22 shall have that license automatically placed into inactive
- 23 status upon issuance of a physician assistant license. Any
- 24 person who holds an active license as a physician assistant
- 25 who is issued a license or permit pursuant to the Medical

- 1 Practice Act of 1987 shall have the physician assistant
- 2 license automatically placed into inactive status.
- 3 Any physician assistant requesting restoration from
- 4 inactive status shall be required to pay the current renewal
- 5 fee and shall be required to restore his or her license, as
- 6 provided in Section 16 of this Act.
- 7 Any physician assistant whose license is in an inactive
- 8 status shall not practice in the State of Illinois.
- 9 Any licensee who shall engage in practice while his or her
- 10 license is lapsed or on inactive status shall be considered to
- 11 be practicing without a license, which shall be grounds for
- discipline under Section 21 of this Act.
- 13 (Source: P.A. 90-61, eff. 12-30-97.)
- 14 (225 ILCS 95/21) (from Ch. 111, par. 4621)
- 15 (Section scheduled to be repealed on January 1, 2028)
- 16 Sec. 21. Grounds for disciplinary action.
- 17 (a) The Department may refuse to issue or to renew, or may
- 18 revoke, suspend, place on probation, reprimand, or take other
- 19 disciplinary or non-disciplinary action with regard to any
- 20 license issued under this Act as the Department may deem
- 21 proper, including the issuance of fines not to exceed \$10,000
- 22 for each violation, for any one or combination of the
- 23 following causes:
- 24 (1) Material misstatement in furnishing information to
- 25 the Department.

- 1 (2) Violations of this Act, or the rules adopted under this Act.
  - (3) Conviction by plea of guilty or nolo contendere, finding of guilt, jury verdict, or entry of judgment or sentencing, including, but not limited to, convictions, preceding sentences of supervision, conditional discharge, or first offender probation, under the laws of any jurisdiction of the United States that is: (i) a felony; or (ii) a misdemeanor, an essential element of which is dishonesty, or that is directly related to the practice of the profession.
  - (4) Making any misrepresentation for the purpose of obtaining licenses.
    - (5) Professional incompetence.
  - (6) Aiding or assisting another person in violating any provision of this Act or its rules.
  - (7) Failing, within 60 days, to provide information in response to a written request made by the Department.
  - (8) Engaging in dishonorable, unethical, or unprofessional conduct, as defined by rule, of a character likely to deceive, defraud, or harm the public.
  - (9) Habitual or excessive use or addiction to alcohol, narcotics, stimulants, or any other chemical agent or drug that results in a physician assistant's inability to practice with reasonable judgment, skill, or safety.
    - (10) Discipline by another U.S. jurisdiction or

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foreign nation, if at least one of the grounds for discipline is the same or substantially equivalent to those set forth in this Section.

- (11) Directly or indirectly giving to or receiving from any person, firm, corporation, partnership, association any fee, commission, rebate or other form of compensation for any professional services not actually or personally rendered. Nothing in this paragraph affects any bona fide independent contractor or employment arrangements, which may include provisions for compensation, health insurance, pension, other or employment benefits, with persons or entities authorized under this Act for the provision of services within the scope of the licensee's practice under this Act.
- (12) A finding by the <del>Disciplinary</del> Board that the licensee, after having his or her license placed on probationary status has violated the terms of probation.
  - (13) Abandonment of a patient.
- (14) Willfully making or filing false records or reports in his or her practice, including but not limited to false records filed with state agencies or departments.
- (15) Willfully failing to report an instance of suspected child abuse or neglect as required by the Abused and Neglected Child Reporting Act.
- (16) Physical illness, or mental illness or impairment that results in the inability to practice the profession

with reasonable judgment, skill, or safety, including, but not limited to, deterioration through the aging process or loss of motor skill.

- (17) Being named as a perpetrator in an indicated report by the Department of Children and Family Services under the Abused and Neglected Child Reporting Act, and upon proof by clear and convincing evidence that the licensee has caused a child to be an abused child or neglected child as defined in the Abused and Neglected Child Reporting Act.
  - (18) (Blank).
- (19) Gross negligence resulting in permanent injury or death of a patient.
- (20) Employment of fraud, deception or any unlawful means in applying for or securing a license as a physician assistant.
- (21) Exceeding the authority delegated to him or her by his or her collaborating physician in a written collaborative agreement when such agreement is required under this Act.
- (22) Immoral conduct in the commission of any act, such as sexual abuse, sexual misconduct, or sexual exploitation related to the licensee's practice.
- (23) Violation of the Health Care Worker Self-Referral Act.
  - (24) Practicing under a false or assumed name, except

as province sy ran	d by law.	provided	l as
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- (25) Making a false or misleading statement regarding his or her skill or the efficacy or value of the medicine, treatment, or remedy prescribed by him or her in the course of treatment.
- (26) Allowing another person to use his or her license to practice.
- (27) Prescribing, selling, administering, distributing, giving, or self-administering a drug classified as a controlled substance for other than medically accepted therapeutic purposes.
- (28) Promotion of the sale of drugs, devices, appliances, or goods provided for a patient in a manner to exploit the patient for financial gain.
- (29) A pattern of practice or other behavior that demonstrates incapacity or incompetence to practice under this Act.
- (30) Violating State or federal laws or regulations relating to controlled substances or other legend drugs or ephedra as defined in the Ephedra Prohibition Act.
- (31) (Blank). Exceeding the prescriptive authority delegated by the collaborating physician or violating the written collaborative agreement delegating that authority.
- (32) (Blank). Practicing without providing to the Department a notice of collaboration or delegation of prescriptive authority.

- 1 (33) Failure to establish and maintain records of 2 patient care and treatment as required by law.
  - (34) Attempting to subvert or cheat on the examination of the National Commission on Certification of Physician Assistants or its successor agency.
  - (35) Willfully or negligently violating the confidentiality between physician assistant and patient, except as required by law.
  - (36) Willfully failing to report an instance of suspected abuse, neglect, financial exploitation, or self-neglect of an eligible adult as defined in and required by the Adult Protective Services Act.
  - (37) Being named as an abuser in a verified report by the Department on Aging under the Adult Protective Services Act and upon proof by clear and convincing evidence that the licensee abused, neglected, or financially exploited an eligible adult as defined in the Adult Protective Services Act.
  - (38) Failure to report to the Department an adverse final action taken against him or her by another licensing jurisdiction of the United States or a foreign state or country, a peer review body, a health care institution, a professional society or association, a governmental agency, a law enforcement agency, or a court acts or conduct similar to acts or conduct that would constitute grounds for action under this Section.

1	(	(39)	Failure	to	provide	е	copies	of	records	of	patient
2	care	or	treatment	, e:	xcept as	s i	required	d b	/ law.		

- (40) (Blank). Entering into an excessive number of written collaborative agreements with licensed physicians resulting in an inability to adequately collaborate.
- (41) (Blank). Repeated failure to adequately collaborate with a collaborating physician.
- (42) Violating the Compassionate Use of Medical Cannabis Program Act.
- (b) The Department may, without a hearing, refuse to issue or renew or may suspend the license of any person who fails to file a return, or to pay the tax, penalty or interest shown in a filed return, or to pay any final assessment of the tax, penalty, or interest as required by any tax Act administered by the Illinois Department of Revenue, until such time as the requirements of any such tax Act are satisfied.
- (c) The determination by a circuit court that a licensee is subject to involuntary admission or judicial admission as provided in the Mental Health and Developmental Disabilities Code operates as an automatic suspension. The suspension will end only upon a finding by a court that the patient is no longer subject to involuntary admission or judicial admission and issues an order so finding and discharging the patient, and upon the recommendation of the Disciplinary Board to the Secretary that the licensee be allowed to resume his or her practice.

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(d) In enforcing this Section, the Department upon a showing of a possible violation may compel an individual licensed to practice under this Act, or who has applied for licensure under this Act, to submit to a mental or physical examination, or both, which may include a substance abuse or sexual offender evaluation, as required by and at the expense of the Department.

The Department shall specifically designate the examining physician licensed to practice medicine in all of its branches or, if applicable, the multidisciplinary team involved in providing the mental or physical examination or both. The multidisciplinary team shall be led by a physician licensed to practice medicine in all of its branches and may consist of one or more or a combination of physicians licensed to practice all medicine in of its branches, licensed clinical psychologists, licensed clinical social workers, licensed clinical professional counselors, and other professional and administrative staff. Any examining physician or member of the multidisciplinary team may require any person ordered to submit to an examination pursuant to this Section to submit to supplemental testing deemed necessary to any additional complete any examination or evaluation process, including, but not limited to, blood testing, urinalysis, psychological testing, or neuropsychological testing.

The Department may order the examining physician or any member of the multidisciplinary team to provide to the

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1 Department any and all records, including business records,

that relate to the examination and evaluation, including any

3 supplemental testing performed.

The Department may order the examining physician or any member of the multidisciplinary team to present testimony concerning the mental or physical examination of the licensee applicant. No information, report, record, or other documents in any way related to the examination shall be excluded by reason of any common law or statutory privilege relating to communications between the licensee or applicant and the examining physician or any member of the multidisciplinary team. No authorization is necessary from the licensee or applicant ordered to undergo an examination for the examining physician or any member of the multidisciplinary team to provide information, reports, records, or other documents or to provide any testimony regarding the examination and evaluation.

The individual to be examined may have, at his or her own expense, another physician of his or her choice present during all aspects of this examination. However, that physician shall be present only to observe and may not interfere in any way with the examination.

Failure of an individual to submit to a mental or physical examination, when ordered, shall result in an automatic suspension of his or her license until the individual submits to the examination.

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If the Department finds an individual unable to practice because of the reasons set forth in this Section, the Department may require that individual to submit to care, counseling, or treatment by physicians approved or designated by the Department, as a condition, term, or restriction for continued, reinstated, or renewed licensure to practice; or, in lieu of care, counseling, or treatment, the Department may file a complaint to immediately suspend, revoke, or otherwise discipline the license of the individual. An individual whose license was granted, continued, reinstated, disciplined, or supervised subject to such terms, conditions, or restrictions, and who fails to comply with such terms, conditions, or restrictions, shall be referred to Secretary for a determination as to whether the individual shall have his or her license suspended immediately, pending a hearing by the Department.

In instances in which the Secretary immediately suspends a person's license under this Section, a hearing on that person's license must be convened by the Department within 30 days after the suspension and completed without appreciable delay. The Department shall have the authority to review the subject individual's record of treatment and counseling regarding the impairment to the extent permitted by applicable federal statutes and regulations safeguarding the confidentiality of medical records.

An individual licensed under this Act and affected under

- this Section shall be afforded an opportunity to demonstrate to the Department that he or she can resume practice in compliance with acceptable and prevailing standards under the provisions of his or her license.
  - (e) An individual or organization acting in good faith, and not in a willful and wanton manner, in complying with this Section by providing a report or other information to the Board, by assisting in the investigation or preparation of a report or information, by participating in proceedings of the Board, or by serving as a member of the Board, shall not be subject to criminal prosecution or civil damages as a result of such actions.
  - (f) Members of the Board and the Disciplinary Board shall be indemnified by the State for any actions occurring within the scope of services on the Disciplinary Board or Board, done in good faith and not willful and wanton in nature. The Attorney General shall defend all such actions unless he or she determines either that there would be a conflict of interest in such representation or that the actions complained of were not in good faith or were willful and wanton.

If the Attorney General declines representation, the member has the right to employ counsel of his or her choice, whose fees shall be provided by the State, after approval by the Attorney General, unless there is a determination by a court that the member's actions were not in good faith or were willful and wanton.

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The member must notify the Attorney General within 7 days after receipt of notice of the initiation of any action involving services of the <del>Disciplinary</del> Board. Failure to so notify the Attorney General constitutes an absolute waiver of the right to a defense and indemnification.

The Attorney General shall determine, within 7 days after receiving such notice, whether he or she will undertake to represent the member.

9 (Source: P.A. 101-363, eff. 8-9-19; 102-558, eff. 8-20-21.)

10 (225 ILCS 95/22.2) (from Ch. 111, par. 4622.2)

(Section scheduled to be repealed on January 1, 2028)

Sec. 22.2. Investigation; notice; hearing. The Department may investigate the actions of any applicant or of any person or persons holding or claiming to hold a license. The Department shall, before suspending, revoking, placing on probationary status, or taking any other disciplinary action as the Department may deem proper with regard to any license, at least 30 days prior to the date set for the hearing, notify the applicant or licensee in writing of any charges made and the time and place for a hearing of the charges before the Disciplinary Board, direct him or her to file his or her written answer thereto to the Disciplinary Board under oath within 20 days after the service on him or her of such notice and inform him or her that if he or she fails to file such answer default will be taken against him or her and his or her

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license may be suspended, revoked, placed on probationary status, or have other disciplinary action, including limiting the scope, nature or extent of his or her practice, as the Department may deem proper taken with regard thereto. Written or electronic notice may be served by personal delivery, email, or mail to the applicant or licensee at his or her address of record or email address of record. At the time and place fixed in the notice, the Department shall proceed to hear the charges and the parties or their counsel shall be accorded ample opportunity to present such statements, testimony, evidence, and argument as may be pertinent to the charges or to the defense thereto. The Department may continue such hearing from time to time. In case the applicant or licensee, after receiving notice, fails to file an answer, his or her license may in the discretion of the Secretary, having received first the recommendation of the Disciplinary Board, be suspended, revoked, placed on probationary status, or the Secretary may take whatever disciplinary action as he or she may deem proper, including limiting the scope, nature, or extent of such person's practice, without a hearing, if the act or acts charged constitute sufficient grounds for such action under this Act.

23 (Source: P.A. 100-453, eff. 8-25-17.)

24 (225 ILCS 95/22.3) (from Ch. 111, par. 4622.3)

(Section scheduled to be repealed on January 1, 2028)

Sec. 22.3. The Department, at its expense, shall preserve 1 a record of all proceedings at the formal hearing of any case 2 involving the refusal to issue, renew or discipline of a 3 license. The notice of hearing, complaint and all other 4 5 documents in the nature of pleadings and written motions filed in the proceedings, the transcript of testimony, the report of 6 the <del>Disciplinary</del> Board or hearing officer and orders of the 7 8 Department shall be the record of such proceeding.

- 9 (Source: P.A. 85-981.)
- 10 (225 ILCS 95/22.5) (from Ch. 111, par. 4622.5)
- 11 (Section scheduled to be repealed on January 1, 2028)
- Sec. 22.5. Subpoena power; oaths. The Department shall have power to subpoena and bring before it any person and to take testimony either orally or by deposition or both, with the same fees and mileage and in the same manner as prescribed by law in judicial proceedings in civil cases in circuit
- 17 courts of this State.

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The Secretary, the designated hearing officer, and any member of the <del>Disciplinary</del> Board designated by the Secretary shall each have power to administer oaths to witnesses at any hearing which the Department is authorized to conduct under this Act and any other oaths required or authorized to be

administered by the Department under this Act.

24 (Source: P.A. 95-703, eff. 12-31-07.)

- 1 (225 ILCS 95/22.6) (from Ch. 111, par. 4622.6)
- 2 (Section scheduled to be repealed on January 1, 2028)
- Sec. 22.6. At the conclusion of the hearing, the

  Disciplinary Board shall present to the Secretary a written

  report of its findings of fact, conclusions of law, and

  recommendations. The report shall contain a finding whether or

  not the accused person violated this Act or failed to comply

  with the conditions required in this Act. The Disciplinary
- 9 Board shall specify the nature of the violation or failure to
  10 comply, and shall make its recommendations to the Secretary.
  11 The report of findings of fact, conclusions of law, and
  12 recommendation of the Disciplinary Board shall be the basis
- 14 license or permit. If the Secretary disagrees in any regard

for the Department's order or refusal or for the granting of a

- with the report of the <del>Disciplinary</del> Board, the Secretary may
- issue an order in contravention thereof. The finding is not
- 17 admissible in evidence against the person in a criminal
- 18 prosecution brought for the violation of this Act, but the
- 19 hearing and finding are not a bar to a criminal prosecution
- 20 brought for the violation of this Act.
- 21 (Source: P.A. 100-453, eff. 8-25-17.)
- 22 (225 ILCS 95/22.7) (from Ch. 111, par. 4622.7)
- 23 (Section scheduled to be repealed on January 1, 2028)
- Sec. 22.7. Hearing officer. Notwithstanding the provisions
- of Section 22.2 of this Act, the Secretary shall have the

authority to appoint any attorney duly licensed to practice 1 2 law in the State of Illinois to serve as the hearing officer in any action for refusal to issue or renew, or for discipline of, 3 a license. The hearing officer shall have full authority to 5 conduct the hearing. The hearing officer shall report his or her findings of fact, conclusions of law, and recommendations 6 to the <del>Disciplinary</del> Board and the Secretary. The <del>Disciplinary</del> 7 8 Board shall have 60 days from receipt of the report to review 9 the report of the hearing officer and present their findings 10 of fact, conclusions of law, and recommendations to the 11 Secretary. If the <del>Disciplinary</del> Board fails to present its 12 report within the 60-day period, the respondent may request in writing a direct appeal to the Secretary, in which case the 13 14 Secretary may issue an order based upon the report of the 15 hearing officer and the record of the proceedings or issue an 16 order remanding the matter back to the hearing officer for 17 proceedings in accordance with the additional Notwithstanding any other provision of this Section, if the 18 Secretary, upon review, determines that substantial justice 19 20 has not been done in the revocation, suspension, or refusal to issue or renew a license or other disciplinary action taken as 21 22 the result of the entry of the hearing officer's report, the 23 Secretary may order a rehearing by the same or other examiners. If the Secretary disagrees in any regard with the 24 25 report of the Disciplinary Board or hearing officer, he or she 26 may issue an order in contravention thereof.

1 (Source: P.A. 100-453, eff. 8-25-17.)

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(225 ILCS 95/22.8) (from Ch. 111, par. 4622.8)
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          (Section scheduled to be repealed on January 1, 2028)
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          Sec. 22.8. In any case involving the refusal to issue,
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      renew or discipline of a license, a copy of the Disciplinary
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      Board's report shall be served upon the respondent by the
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      Department, either personally or as provided in this Act for
      the service of the notice of hearing. Within 20 days after such
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      service, the respondent may present to the Department a motion
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      in writing for a rehearing, which motion shall specify the
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      particular grounds therefor. If no motion for rehearing is
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      filed, then upon the expiration of the time specified for
      filing such a motion, or if a motion for rehearing is denied,
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      then upon such denial the Secretary may enter an order in
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      accordance with recommendations of the Disciplinary Board
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      except as provided in Section 22.6 or 22.7 of this Act. If the
      respondent shall order from the reporting service, and pay for
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      a transcript of the record within the time for filing a motion
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      for rehearing, the 20 day period within which such a motion may
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      be filed shall commence upon the delivery of the transcript to
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      the respondent.
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22 (Source: P.A. 95-703, eff. 12-31-07.)

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23 (225 ILCS 95/22.9) (from Ch. 111, par. 4622.9)

(Section scheduled to be repealed on January 1, 2028)

- 1 Sec. 22.9. Whenever the Secretary is satisfied that
- 2 substantial justice has not been done in the revocation,
- 3 suspension or refusal to issue or renew a license, the
- 4 Secretary may order a rehearing by the same or another hearing
- 5 officer or <del>Disciplinary</del> Board.
- 6 (Source: P.A. 95-703, eff. 12-31-07.)
- 7 (225 ILCS 95/22.10) (from Ch. 111, par. 4622.10)
- 8 (Section scheduled to be repealed on January 1, 2028)
- 9 Sec. 22.10. Order or certified copy; prima facie proof. An
- 10 order or a certified copy thereof, over the seal of the
- 11 Department and purporting to be signed by the Secretary, shall
- 12 be prima facie proof that:
- 13 (a) the signature is the genuine signature of the
- 14 Secretary;
- 15 (b) the Secretary is duly appointed and qualified; and
- 16 (c) the <del>Disciplinary</del> Board and the members thereof are
- 17 qualified to act.
- 18 (Source: P.A. 95-703, eff. 12-31-07.)
- 19 Section 10. The Illinois Controlled Substances Act is
- amended by changing Sections 102 and 303.05 as follows:
- 21 (720 ILCS 570/102) (from Ch. 56 1/2, par. 1102)
- Sec. 102. Definitions. As used in this Act, unless the
- 23 context otherwise requires:

- (a) "Addict" means any person who habitually uses any drug, chemical, substance or dangerous drug other than alcohol so as to endanger the public morals, health, safety or welfare or who is so far addicted to the use of a dangerous drug or controlled substance other than alcohol as to have lost the power of self control with reference to his or her addiction.
  - (b) "Administer" means the direct application of a controlled substance, whether by injection, inhalation, ingestion, or any other means, to the body of a patient, research subject, or animal (as defined by the Humane Euthanasia in Animal Shelters Act) by:
- 12 (1) a practitioner (or, in his or her presence, by his or her authorized agent),
- 14 (2) the patient or research subject pursuant to an order, or
  - (3) a euthanasia technician as defined by the Humane Euthanasia in Animal Shelters Act.
  - (c) "Agent" means an authorized person who acts on behalf of or at the direction of a manufacturer, distributor, dispenser, prescriber, or practitioner. It does not include a common or contract carrier, public warehouseman or employee of the carrier or warehouseman.
    - (c-1) "Anabolic Steroids" means any drug or hormonal substance, chemically and pharmacologically related to testosterone (other than estrogens, progestins, corticosteroids, and dehydroepiandrosterone), and includes:

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(i) 3[beta], 17-dihydroxy-5a-androstane,
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           (ii) 3[alpha], 17[beta]-dihydroxy-5a-androstane,
           (iii) 5[alpha]-androstan-3,17-dione,
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           (iv) 1-androstenediol (3[beta],
 4
              17[beta]-dihydroxy-5[alpha]-androst-1-ene),
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           (v) 1-androstenediol (3[alpha],
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              17[beta]-dihydroxy-5[alpha]-androst-1-ene),
           (vi) 4-androstenediol
 8
 9
               (3[beta], 17[beta]-dihydroxy-androst-4-ene),
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           (vii) 5-androstenediol
11
               (3[beta], 17[beta]-dihydroxy-androst-5-ene),
12
           (viii) 1-androstenedione
13
               ([5alpha]-androst-1-en-3,17-dione),
           (ix) 4-androstenedione
14
               (androst-4-en-3,17-dione),
15
16
           (x) 5-androstenedione
17
               (androst-5-en-3,17-dione),
           (xi) bolasterone (7[alpha], 17a-dimethyl-17[beta]-
18
19
              hydroxyandrost-4-en-3-one),
20
           (xii) boldenone (17[beta]-hydroxyandrost-
              1,4,-diene-3-one),
21
22
           (xiii) boldione (androsta-1,4-
23
              diene-3,17-dione),
24
           (xiv) calusterone (7[beta], 17[alpha]-dimethyl-17
25
               [beta]-hydroxyandrost-4-en-3-one),
           (xv) clostebol (4-chloro-17[beta]-
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hydroxyandrost-4-en-3-one),
1
 2
          (xvi) dehydrochloromethyltestosterone (4-chloro-
              17[beta]-hydroxy-17[alpha]-methyl-
 3
              androst-1, 4-dien-3-one),
 4
 5
          (xvii) desoxymethyltestosterone
 6
          (17[alpha]-methyl-5[alpha]
              -androst-2-en-17[beta]-ol)(a.k.a., madol),
7
 8
          (xviii) [delta]1-dihydrotestosterone (a.k.a.
 9
               '1-testosterone') (17[beta]-hydroxy-
10
              5[alpha]-androst-1-en-3-one),
11
          (xix) 4-dihydrotestosterone (17[beta]-hydroxy-
12
              androstan-3-one),
13
          (xx) drostanolone (17[beta]-hydroxy-2[alpha]-methyl-
14
              5[alpha]-androstan-3-one),
15
          (xxi) ethylestrenol (17[alpha]-ethyl-17[beta]-
16
              hydroxyestr-4-ene),
17
          (xxii) fluoxymesterone (9-fluoro-17[alpha]-methyl-
              1[beta], 17[beta]-dihydroxyandrost-4-en-3-one),
18
          (xxiii) formebolone (2-formyl-17[alpha]-methyl-11[alpha],
19
20
              17[beta]-dihydroxyandrost-1,4-dien-3-one),
21
          (xxiv) furazabol (17[alpha]-methyl-17[beta]-
22
              hydroxyandrostano[2,3-c]-furazan),
23
          (xxv) 13[beta]-ethyl-17[beta]-hydroxygon-4-en-3-one,
          (xxvi) 4-hydroxytestosterone (4,17[beta]-dihydroxy-
24
25
              androst-4-en-3-one),
26
          (xxvii) 4-hydroxy-19-nortestosterone (4,17[beta]-
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dihydroxy-estr-4-en-3-one),
1
 2
          (xxviii) mestanolone (17[alpha]-methyl-17[beta]-
              hydroxy-5-androstan-3-one),
 3
 4
          (xxix) mesterolone (1amethyl-17[beta]-hydroxy-
 5
              [5a]-androstan-3-one),
          (xxx) methandienone (17[alpha]-methyl-17[beta]-
 6
              hydroxyandrost-1,4-dien-3-one),
7
          (xxxi) methandriol (17[alpha]-methyl-3[beta],17[beta]-
 8
 9
              dihydroxyandrost-5-ene),
10
          (xxxii) methenolone (1-methyl-17[beta]-hydroxy-
11
              5[alpha]-androst-1-en-3-one),
12
          (xxxiii) 17[alpha]-methyl-3[beta], 17[beta]-
13
              dihydroxy-5a-androstane,
          (xxxiv) 17[alpha]-methyl-3[alpha],17[beta]-dihydroxy
14
15
              -5a-androstane.
16
          (xxxv) 17[alpha]-methyl-3[beta],17[beta]-
17
              dihydroxyandrost-4-ene),
          (xxxvi) 17[alpha]-methyl-4-hydroxynandrolone (17[alpha]-
18
              methyl-4-hydroxy-17[beta]-hydroxyestr-4-en-3-one),
19
20
          (xxxvii) methyldienolone (17[alpha]-methyl-17[beta]-
              hydroxyestra-4,9(10)-dien-3-one),
21
22
          (xxxviii) methyltrienolone (17[alpha]-methyl-17[beta]-
23
              hydroxyestra-4,9-11-trien-3-one),
          (xxxix) methyltestosterone (17[alpha]-methyl-17[beta]-
24
25
              hydroxyandrost-4-en-3-one),
26
          (x1) mibolerone (7[alpha],17a-dimethyl-17[beta]-
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hydroxyestr-4-en-3-one),
1
 2
          (xli) 17[alpha]-methyl-[delta]1-dihydrotestosterone
 3
               (17b[beta]-hydroxy-17[alpha]-methyl-5[alpha]-
 4
              androst-1-en-3-one) (a.k.a. '17-[alpha]-methyl-
 5
              1-testosterone'),
          (xlii) nandrolone (17[beta]-hydroxyestr-4-en-3-one),
 6
          (xliii) 19-nor-4-androstenediol (3[beta], 17[beta]-
7
 8
              dihydroxyestr-4-ene),
 9
          (xliv) 19-nor-4-androstenediol (3[alpha], 17[beta]-
              dihydroxyestr-4-ene),
10
11
          (xlv) 19-nor-5-androstenediol (3[beta], 17[beta]-
12
              dihydroxyestr-5-ene),
13
          (xlvi) 19-nor-5-androstenediol (3[alpha], 17[beta]-
14
              dihydroxyestr-5-ene),
15
          (xlvii) 19-nor-4,9(10)-androstadienedione
16
               (estra-4,9(10)-diene-3,17-dione),
          (xlviii) 19-nor-4-androstenedione (estr-4-
17
              en-3,17-dione),
18
          (xlix) 19-nor-5-androstenedione (estr-5-
19
20
              en-3,17-dione),
          (1) norbolethone (13[beta], 17a-diethyl-17[beta]-
21
22
              hydroxygon-4-en-3-one),
23
          (li) norclostebol (4-chloro-17[beta]-
              hydroxyestr-4-en-3-one),
24
25
          (lii) norethandrolone (17[alpha]-ethyl-17[beta]-
26
              hydroxyestr-4-en-3-one),
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(liii) normethandrolone (17[alpha]-methyl-17[beta]-
1
              hydroxyestr-4-en-3-one),
 2
          (liv) oxandrolone (17[alpha]-methyl-17[beta]-hydroxy-
 3
 4
              2-oxa-5[alpha]-androstan-3-one),
 5
          (lv) oxymesterone (17[alpha]-methyl-4,17[beta]-
              dihydroxyandrost-4-en-3-one),
 6
          (lvi) oxymetholone (17[alpha]-methyl-2-hydroxymethylene-
7
              17[beta]-hydroxy-(5[alpha]-androstan-3-one),
 8
 9
          (lvii) stanozolol (17[alpha]-methyl-17[beta]-hydroxy-
10
              (5[alpha]-androst-2-eno[3,2-c]-pyrazole),
11
          (lviii) stenbolone (17[beta]-hydroxy-2-methyl-
12
              (5[alpha]-androst-1-en-3-one),
13
          (lix) testolactone (13-hydroxy-3-oxo-13,17-
              secoandrosta-1,4-dien-17-oic
14
15
              acid lactone),
16
          (lx) testosterone (17[beta]-hydroxyandrost-
17
              4-en-3-one),
          (lxi) tetrahydrogestrinone (13[beta], 17[alpha]-
18
19
              diethyl-17[beta]-hydroxygon-
20
              4,9,11-trien-3-one),
          (lxii) trenbolone (17[beta]-hydroxyestr-4,9,
21
22
              11-trien-3-one).
23
          Any person who is otherwise lawfully in possession of an
      anabolic steroid, or who otherwise lawfully manufactures,
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      distributes, dispenses, delivers, or possesses with intent to
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      deliver an anabolic steroid, which anabolic steroid is
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- expressly intended for and lawfully allowed to be administered through implants to livestock or other nonhuman species, and which is approved by the Secretary of Health and Human Services for such administration, and which the person intends to administer or have administered through such implants, shall not be considered to be in unauthorized possession or to unlawfully manufacture, distribute, dispense, deliver, or possess with intent to deliver such anabolic steroid for purposes of this Act.
- (d) "Administration" means the Drug Enforcement Administration, United States Department of Justice, or its successor agency.
  - (d-5) "Clinical Director, Prescription Monitoring Program" means a Department of Human Services administrative employee licensed to either prescribe or dispense controlled substances who shall run the clinical aspects of the Department of Human Services Prescription Monitoring Program and its Prescription Information Library.
  - (d-10) "Compounding" means the preparation and mixing of components, excluding flavorings, (1) as the result of a prescriber's prescription drug order or initiative based on the prescriber-patient-pharmacist relationship in the course of professional practice or (2) for the purpose of, or incident to, research, teaching, or chemical analysis and not for sale or dispensing. "Compounding" includes the preparation of drugs or devices in anticipation of receiving prescription

- drug orders based on routine, regularly observed dispensing patterns. Commercially available products may be compounded for dispensing to individual patients only if both of the following conditions are met: (i) the commercial product is not reasonably available from normal distribution channels in a timely manner to meet the patient's needs and (ii) the prescribing practitioner has requested that the drug be compounded.
  - (e) "Control" means to add a drug or other substance, or immediate precursor, to a Schedule whether by transfer from another Schedule or otherwise.
  - (f) "Controlled Substance" means (i) a drug, substance, immediate precursor, or synthetic drug in the Schedules of Article II of this Act or (ii) a drug or other substance, or immediate precursor, designated as a controlled substance by the Department through administrative rule. The term does not include distilled spirits, wine, malt beverages, or tobacco, as those terms are defined or used in the Liquor Control Act of 1934 and the Tobacco Products Tax Act of 1995.
  - (f-5) "Controlled substance analog" means a substance:
    - (1) the chemical structure of which is substantially similar to the chemical structure of a controlled substance in Schedule I or II;
      - (2) which has a stimulant, depressant, or hallucinogenic effect on the central nervous system that is substantially similar to or greater than the stimulant,

- depressant, or hallucinogenic effect on the central nervous system of a controlled substance in Schedule I or II; or
  - (3) with respect to a particular person, which such person represents or intends to have a stimulant, depressant, or hallucinogenic effect on the central nervous system that is substantially similar to or greater than the stimulant, depressant, or hallucinogenic effect on the central nervous system of a controlled substance in Schedule I or II.
  - (g) "Counterfeit substance" means 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.
  - (h) "Deliver" or "delivery" means the actual, constructive or attempted transfer of possession of a controlled substance, with or without consideration, whether or not there is an agency relationship. "Deliver" or "delivery" does not include the donation of drugs to the extent permitted under the Illinois Drug Reuse Opportunity Program Act.
  - (i) "Department" means the Illinois Department of Human Services (as successor to the Department of Alcoholism and Substance Abuse) or its successor agency.

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- 1 (j) (Blank).
- 2 (k) "Department of Corrections" means the Department of Corrections of the State of Illinois or its successor agency.
- 4 (1) "Department of Financial and Professional Regulation"
  5 means the Department of Financial and Professional Regulation
  6 of the State of Illinois or its successor agency.
  - (m) "Depressant" means any drug that (i) causes an overall depression of central nervous system functions, (ii) causes impaired consciousness and awareness, and (iii) can be habit-forming or lead to a substance abuse problem, including, but not limited to, alcohol, cannabis and its active principles and their analogs, benzodiazepines and their analogs, barbiturates and their analogs, opioids (natural and synthetic) and their analogs, and chloral hydrate and similar sedative hypnotics.
- 16 (n) (Blank).
- 17 (o) "Director" means the Director of the Illinois State
  18 Police or his or her designated agents.
- 19 (p) "Dispense" means to deliver a controlled substance to 20 an ultimate user or research subject by or pursuant to the 21 lawful order of a prescriber, including the prescribing, 22 administering, packaging, labeling, or compounding necessary 23 to prepare the substance for that delivery.
- 24 (q) "Dispenser" means a practitioner who dispenses.
- 25 (r) "Distribute" means to deliver, other than by administering or dispensing, a controlled substance.

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- 1 (s) "Distributor" means a person who distributes.
- 2 (t) "Drug" means (1) substances recognized as drugs in the 3 official United States Pharmacopoeia, Official Homeopathic Pharmacopoeia of the United States, or official National 5 Formulary, or any supplement to any of them; (2) substances intended for use in diagnosis, cure, mitigation, treatment, or 6 7 prevention of disease in man or animals; (3) substances (other than food) intended to affect the structure of any function of 8 9 the body of man or animals and (4) substances intended for use 10 as a component of any article specified in clause (1), (2), or 11 (3) of this subsection. It does not include devices or their 12 components, parts, or accessories.
  - (t-3) "Electronic health record" or "EHR" means an electronic record of health-related information on an individual that is created, gathered, managed, and consulted by authorized health care clinicians and staff.
  - means any computer-based system or combination of federally certified Health IT Modules (defined at 42 CFR 170.102 or its successor) used as a repository for electronic health records and accessed or updated by a prescriber or authorized surrogate in the ordinary course of his or her medical practice. For purposes of connecting to the Prescription Information Library maintained by the Bureau of Pharmacy and Clinical Support Systems or its successor, an EHR system may connect to the Prescription Information Library directly or

- 1 through all or part of a computer program or system that is a
- 2 federally certified Health IT Module maintained by a third
- 3 party and used by the EHR system to secure access to the
- 4 database.
- 5 (t-4) "Emergency medical services personnel" has the
- 6 meaning ascribed to it in the Emergency Medical Services (EMS)
- 7 Systems Act.
- 8 (t-5) "Euthanasia agency" means an entity certified by the
- 9 Department of Financial and Professional Regulation for the
- 10 purpose of animal euthanasia that holds an animal control
- 11 facility license or animal shelter license under the Animal
- 12 Welfare Act. A euthanasia agency is authorized to purchase,
- 13 store, possess, and utilize Schedule II nonnarcotic and
- 14 Schedule III nonnarcotic drugs for the sole purpose of animal
- 15 euthanasia.
- 16 (t-10) "Euthanasia drugs" means Schedule II or Schedule
- 17 III substances (nonnarcotic controlled substances) that are
- 18 used by a euthanasia agency for the purpose of animal
- 19 euthanasia.
- 20 (u) "Good faith" means the prescribing or dispensing of a
- 21 controlled substance by a practitioner in the regular course
- of professional treatment to or for any person who is under his
- or her treatment for a pathology or condition other than that
- 24 individual's physical or psychological dependence upon or
- 25 addiction to a controlled substance, except as provided
- herein: and application of the term to a pharmacist shall mean

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- the dispensing of a controlled substance pursuant to the prescriber's order which in the professional judgment of the pharmacist is lawful. The pharmacist shall be guided by accepted professional standards, including, but not limited
- 6 (1) lack of consistency of prescriber-patient

to, the following, in making the judgment:

relationship,

- (2) frequency of prescriptions for same drug by one prescriber for large numbers of patients,
  - (3) quantities beyond those normally prescribed,
- 11 (4) unusual dosages (recognizing that there may be
  12 clinical circumstances where more or less than the usual
  13 dose may be used legitimately),
- 14 (5) unusual geographic distances between patient,
  15 pharmacist and prescriber,
  - (6) consistent prescribing of habit-forming drugs.
- 17 (u-0.5) "Hallucinogen" means a drug that causes markedly
  18 altered sensory perception leading to hallucinations of any
  19 type.
- 20 (u-1) "Home infusion services" means services provided by
  21 a pharmacy in compounding solutions for direct administration
  22 to a patient in a private residence, long-term care facility,
  23 or hospice setting by means of parenteral, intravenous,
  24 intramuscular, subcutaneous, or intraspinal infusion.
- 25 (u-5) "Illinois State Police" means the Illinois State
  26 Police or its successor agency.

- 1 (v) "Immediate precursor" means a substance:
  - (1) which the Department has found to be and by rule designated as being a principal compound used, or produced primarily for use, in the manufacture of a controlled substance;
    - (2) which is an immediate chemical intermediary used or likely to be used in the manufacture of such controlled substance; and
    - (3) the control of which is necessary to prevent, curtail or limit the manufacture of such controlled substance.
  - (w) "Instructional activities" means the acts of teaching, educating or instructing by practitioners using controlled substances within educational facilities approved by the State Board of Education or its successor agency.
  - (x) "Local authorities" means a duly organized State, County or Municipal peace unit or police force.
  - (y) "Look-alike substance" means a substance, other than a controlled substance which (1) by overall dosage unit appearance, including shape, color, size, markings or lack thereof, taste, consistency, or any other identifying physical characteristic of the substance, would lead a reasonable person to believe that the substance is a controlled substance, or (2) is expressly or impliedly represented to be a controlled substance or is distributed under circumstances which would lead a reasonable person to believe that the

- substance is a controlled substance. For the purpose of determining whether the representations made or the circumstances of the distribution would lead a reasonable person to believe the substance to be a controlled substance under this clause (2) of subsection (y), the court or other authority may consider the following factors in addition to any other factor that may be relevant:
  - (a) statements made by the owner or person in control of the substance concerning its nature, use or effect;
  - (b) statements made to the buyer or recipient that the substance may be resold for profit;
  - (c) whether the substance is packaged in a manner normally used for the illegal distribution of controlled substances;
  - (d) whether the distribution or attempted distribution included an exchange of or demand for money or other property as consideration, and whether the amount of the consideration was substantially greater than the reasonable retail market value of the substance.
- Clause (1) of this subsection (y) shall not apply to a noncontrolled substance in its finished dosage form that was initially introduced into commerce prior to the initial introduction into commerce of a controlled substance in its finished dosage form which it may substantially resemble.
- Nothing in this subsection (y) prohibits the dispensing or distributing of noncontrolled substances by persons authorized

- 1 to dispense and distribute controlled substances under this
- 2 Act, provided that such action would be deemed to be carried
- 3 out in good faith under subsection (u) if the substances
- 4 involved were controlled substances.
- 5 Nothing in this subsection (y) or in this Act prohibits
- 6 the manufacture, preparation, propagation, compounding,
- 7 processing, packaging, advertising or distribution of a drug
- 8 or drugs by any person registered pursuant to Section 510 of
- 9 the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360).
- 10 (y-1) "Mail-order pharmacy" means a pharmacy that is
- 11 located in a state of the United States that delivers,
- dispenses or distributes, through the United States Postal
- 13 Service or other common carrier, to Illinois residents, any
- substance which requires a prescription.
- 15 (z) "Manufacture" means the production, preparation,
- 16 propagation, compounding, conversion or processing of a
- 17 controlled substance other than methamphetamine, either
- 18 directly or indirectly, by extraction from substances of
- 19 natural origin, or independently by means of chemical
- 20 synthesis, or by a combination of extraction and chemical
- 21 synthesis, and includes any packaging or repackaging of the
- 22 substance or labeling of its container, except that this term
- 23 does not include:
- 24 (1) by an ultimate user, the preparation or
- compounding of a controlled substance for his or her own
- 26 use;

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1	(2)	by a	pract	titione	r, or	his or h	er au	tho	rized agent
2	under	his	or	her	super	vision,	the	р	reparation,
3	compoun	ding,	pac	kaging,	or	labeling	of	a	controlled
4	substan	.ce:							

- (a) as an incident to his or her administering or dispensing of a controlled substance in the course of his or her professional practice; or
- (b) as an incident to lawful research, teaching or chemical analysis and not for sale; or
- (3) the packaging, repackaging, or labeling of drugs only to the extent permitted under the Illinois Drug Reuse Opportunity Program Act.
- 13 (z-1) (Blank).
- 14 (z-5) "Medication shopping" means the conduct prohibited 15 under subsection (a) of Section 314.5 of this Act.
- 16 (z-10) "Mid-level practitioner" means (i) a physician 17 assistant who has been delegated authority to prescribe through a written delegation of authority by a physician 18 19 licensed to practice medicine in all of its branches, in 20 accordance with Section 7.5 of the Physician Assistant Practice Act of 1987, (ii) an advanced practice registered 21 22 nurse who has been delegated authority to prescribe through a 23 written delegation of authority by a physician licensed to practice medicine in all of its branches or by a podiatric 24 25 physician, in accordance with Section 65-40 of the Nurse 26 Practice Act, (iii) an advanced practice registered nurse

- 1 certified as a nurse practitioner, nurse midwife, or clinical
- 2 nurse specialist who has been granted authority to prescribe
- 3 by a hospital affiliate in accordance with Section 65-45 of
- 4 the Nurse Practice Act, (iv) an animal euthanasia agency, or
- 5 (v) a prescribing psychologist.
- 6 (aa) "Narcotic drug" means any of the following, whether
- 7 produced directly or indirectly by extraction from substances
- 8 of vegetable origin, or independently by means of chemical
- 9 synthesis, or by a combination of extraction and chemical
- 10 synthesis:
- 11 (1) opium, opiates, derivatives of opium and opiates,
- including their isomers, esters, ethers, salts, and salts
- of isomers, esters, and ethers, whenever the existence of
- 14 such isomers, esters, ethers, and salts is possible within
- 15 the specific chemical designation; however the term
- 16 "narcotic drug" does not include the isoquinoline
- 17 alkaloids of opium;
- 18 (2) (blank);
- 19 (3) opium poppy and poppy straw;
- 20 (4) coca leaves, except coca leaves and extracts of
- 21 coca leaves from which substantially all of the cocaine
- and ecgonine, and their isomers, derivatives and salts,
- have been removed;
- 24 (5) cocaine, its salts, optical and geometric isomers,
- and salts of isomers;
- 26 (6) ecgonine, its derivatives, their salts, isomers,

- 1 and salts of isomers;
- 2 (7) any compound, mixture, or preparation which
- 3 contains any quantity of any of the substances referred to
- 4 in subparagraphs (1) through (6).
- 5 (bb) "Nurse" means a registered nurse licensed under the
- 6 Nurse Practice Act.
- 7 (cc) (Blank).
- 8 (dd) "Opiate" means any substance having an addiction
- 9 forming or addiction sustaining liability similar to morphine
- or being capable of conversion into a drug having addiction
- 11 forming or addiction sustaining liability.
- 12 (ee) "Opium poppy" means the plant of the species Papaver
- 13 somniferum L., except its seeds.
- 14 (ee-5) "Oral dosage" means a tablet, capsule, elixir, or
- 15 solution or other liquid form of medication intended for
- 16 administration by mouth, but the term does not include a form
- of medication intended for buccal, sublingual, or transmucosal
- 18 administration.
- 19 (ff) "Parole and Pardon Board" means the Parole and Pardon
- 20 Board of the State of Illinois or its successor agency.
- 21 (qq) "Person" means any individual, corporation,
- 22 mail-order pharmacy, government or governmental subdivision or
- 23 agency, business trust, estate, trust, partnership or
- association, or any other entity.
- 25 (hh) "Pharmacist" means any person who holds a license or
- 26 certificate of registration as a registered pharmacist, a

- 1 local registered pharmacist or a registered assistant
- 2 pharmacist under the Pharmacy Practice Act.
- 3 (ii) "Pharmacy" means any store, ship or other place in
- 4 which pharmacy is authorized to be practiced under the
- 5 Pharmacy Practice Act.
- 6 (ii-5) "Pharmacy shopping" means the conduct prohibited
- 7 under subsection (b) of Section 314.5 of this Act.
- 8 (ii-10) "Physician" (except when the context otherwise
- 9 requires) means a person licensed to practice medicine in all
- 10 of its branches.
- 11 (jj) "Poppy straw" means all parts, except the seeds, of
- 12 the opium poppy, after mowing.
- 13 (kk) "Practitioner" means a physician licensed to practice
- 14 medicine in all its branches, dentist, optometrist, podiatric
- 15 physician, veterinarian, scientific investigator, pharmacist,
- 16 physician assistant, advanced practice registered nurse,
- 17 licensed practical nurse, registered nurse, emergency medical
- 18 services personnel, hospital, laboratory, or pharmacy, or
- 19 other person licensed, registered, or otherwise lawfully
- 20 permitted by the United States or this State to distribute,
- 21 dispense, conduct research with respect to, administer or use
- in teaching or chemical analysis, a controlled substance in
- 23 the course of professional practice or research.
- 24 (11) "Pre-printed prescription" means a written
- 25 prescription upon which the designated drug has been indicated
- 26 prior to the time of issuance; the term does not mean a written

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prescription that is individually generated by machine or computer in the prescriber's office.

(mm) "Prescriber" means a physician licensed to practice dentist, optometrist, medicine in all its branches, prescribing psychologist licensed under Section 4.2 of the Psychologist Licensing Act with prescriptive authority delegated under Section 4.3 of the Clinical Psychologist Licensing Act, podiatric physician, veterinarian who issues a prescription, a physician assistant who issues a prescription for a controlled substance in accordance with Section 303.05, a written delegation, and a written collaborative agreement required under Section 7.5 of the Physician Assistant Practice Act of 1987, an advanced practice registered nurse with prescriptive authority delegated under Section 65-40 of the Nurse Practice Act and in accordance with Section 303.05, a written delegation, and a written collaborative agreement under Section 65-35 of the Nurse Practice Act, an advanced practice registered nurse certified as a nurse practitioner, nurse midwife, or clinical nurse specialist who has been granted authority to prescribe by a hospital affiliate in accordance with Section 65-45 of the Nurse Practice Act and in accordance with Section 303.05, or an advanced practice registered nurse certified as a nurse practitioner, nurse midwife, or clinical nurse specialist who has full practice authority pursuant to Section 65-43 of the Nurse Practice Act.

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(nn) "Prescription" means a written, facsimile, or oral order, or an electronic order that complies with applicable federal requirements, of a physician licensed to practice medicine in all its branches, dentist, podiatric physician or veterinarian for any controlled substance, of an optometrist in accordance with Section 15.1 of the Illinois Optometric Practice Act of 1987, of a prescribing psychologist licensed under Section 4.2 of the Clinical Psychologist Licensing Act with prescriptive authority delegated under Section 4.3 of the Clinical Psychologist Licensing Act, of a physician assistant for a controlled substance in accordance with Section 303.05, a written delegation, and a written collaborative agreement required under Section 7.5 of the Physician Assistant Practice Act of 1987, of an advanced practice registered nurse with prescriptive authority delegated under Section 65-40 of the Nurse Practice Act who issues a prescription for a controlled substance in accordance with Section 303.05, a written delegation, and a written collaborative agreement under Section 65-35 of the Nurse Practice Act, of an advanced practice registered nurse certified as a nurse practitioner, nurse midwife, or clinical nurse specialist who has been granted authority to prescribe by a hospital affiliate in accordance with Section 65-45 of the Nurse Practice Act and in accordance with Section 303.05 when required by law, or of an advanced practice registered nurse certified as a nurse practitioner, nurse midwife, or clinical nurse specialist who

- 1 has full practice authority pursuant to Section 65-43 of the
- 2 Nurse Practice Act.
- 3 (nn-5) "Prescription Information Library" (PIL) means an
- 4 electronic library that contains reported controlled substance
- 5 data.
- 6 (nn-10) "Prescription Monitoring Program" (PMP) means the
- 7 entity that collects, tracks, and stores reported data on
- 8 controlled substances and select drugs pursuant to Section
- 9 316.
- 10 (oo) "Production" or "produce" means manufacture,
- 11 planting, cultivating, growing, or harvesting of a controlled
- 12 substance other than methamphetamine.
- 13 (pp) "Registrant" means every person who is required to
- register under Section 302 of this Act.
- 15 (qq) "Registry number" means the number assigned to each
- 16 person authorized to handle controlled substances under the
- 17 laws of the United States and of this State.
- 18 (qq-5) "Secretary" means, as the context requires, either
- 19 the Secretary of the Department or the Secretary of the
- 20 Department of Financial and Professional Regulation, and the
- 21 Secretary's designated agents.
- (rr) "State" includes the State of Illinois and any state,
- 23 district, commonwealth, territory, insular possession thereof,
- 24 and any area subject to the legal authority of the United
- 25 States of America.
- 26 (rr-5) "Stimulant" means any drug that (i) causes an

- 1 overall excitation of central nervous system functions, (ii)
- 2 causes impaired consciousness and awareness, and (iii) can be
- 3 habit-forming or lead to a substance abuse problem, including,
- 4 but not limited to, amphetamines and their analogs,
- 5 methylphenidate and its analogs, cocaine, and phencyclidine
- 6 and its analogs.
- 7 (rr-10) "Synthetic drug" includes, but is not limited to,
- 8 any synthetic cannabinoids or piperazines or any synthetic
- 9 cathinones as provided for in Schedule I.
- 10 (ss) "Ultimate user" means a person who lawfully possesses
- 11 a controlled substance for his or her own use or for the use of
- 12 a member of his or her household or for administering to an
- animal owned by him or her or by a member of his or her
- 14 household.
- 15 (Source: P.A. 101-666, eff. 1-1-22; 102-389, eff. 1-1-22;
- 16 102-538, eff. 8-20-21; 102-813, eff. 5-13-22.)
- 17 (720 ILCS 570/303.05)
- 18 Sec. 303.05. Mid-level practitioner registration.
- 19 (a) The Department of Financial and Professional
- 20 Regulation shall register licensed physician assistants,
- 21 licensed advanced practice registered nurses, and prescribing
- 22 psychologists licensed under Section 4.2 of the Clinical
- 23 Psychologist Licensing Act to prescribe and dispense
- 24 controlled substances under Section 303 and euthanasia
- 25 agencies to purchase, store, or administer animal euthanasia

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- (1) with respect to physician assistants,
- (A) the physician assistant has been delegated written authority to prescribe any Schedule III through V controlled substances by a physician licensed to practice medicine in all its branches in accordance with Section 7.5 of the Physician Assistant Practice Act of 1987; and the physician assistant has completed the appropriate application forms and has paid the required fees as set by rule; or
- (B) the physician assistant has been delegated authority by a collaborating physician licensed to practice medicine in all its branches to prescribe or dispense Schedule II controlled substances through a written delegation of authority and under the following conditions:
  - by oral dosage or topical or transdermal application may be delegated, provided that the delegated Schedule II controlled substances are routinely prescribed by the collaborating physician. This delegation must identify the specific Schedule II controlled substances by either brand name or generic name. Schedule II controlled substances or other route of administration may not be

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1	delegated;
2	(ii) any delegation must be of controlled
3	substances prescribed by the collaborating
4	physician;
5	(iii) all prescriptions must be limited to no
6	more than a 30 day supply, with any continuation
7	authorized only after prior approval of the
8	collaborating physician;
9	(iv) the physician assistant must discuss the
10	condition of any patients for whom a controlled
11	substance is prescribed monthly with the
12	delegating physician;
13	$\underline{\text{(A)}}$ $\underline{\text{(v)}}$ the physician assistant must have
14	completed the appropriate application forms and paid
15	the required fees as set by rule;
16	(B) (vi) the physician assistant must provide
17	evidence of satisfactory completion of 45 contact
18	hours in pharmacology from any physician assistant
19	program accredited by the Accreditation Review
20	Commission on Education for the Physician Assistant
21	(ARC-PA), or its predecessor agency, for any new
22	license issued with Schedule II authority after the

 $\underline{\text{(C)}}$  (vii) the physician assistant must annually complete at least 5 hours of continuing education in

effective date of this amendatory Act of the 97th

General Assembly; and

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- (2) with respect to advanced practice registered nurses who do not meet the requirements of Section 65-43 of the Nurse Practice Act,
  - (A) the advanced practice registered nurse has been delegated authority to prescribe any Schedule III through V controlled substances by a collaborating physician licensed to practice medicine in all its branches or a collaborating podiatric physician in accordance with Section 65-40 of the Nurse Practice Act. The advanced practice registered nurse has completed the appropriate application forms and has paid the required fees as set by rule; or
  - (B) the advanced practice registered nurse has been delegated authority by a collaborating physician licensed to practice medicine in all its branches to prescribe or dispense Schedule II controlled substances through a written delegation of authority and under the following conditions:
    - (i) specific Schedule II controlled substances by oral dosage or topical or transdermal application may be delegated, provided that the delegated Schedule II controlled substances are routinely prescribed by the collaborating physician. This delegation must identify the specific Schedule II controlled substances by

either brand name or generic name. Schedule II 1 2 controlled substances to be delivered by injection 3 other route of administration may not be delegated; (ii) any delegation must be of controlled 6 substances prescribed by the collaborating 7 physician; (iii) all prescriptions must be limited to no 8 9 more than a 30-day supply, with any continuation 10 authorized only after prior approval of the 11 collaborating physician; 12 (iv) the advanced practice registered nurse must discuss the condition of any patients for 13 14 whom a controlled substance is prescribed monthly 15 with the delegating physician or in the course of 16 review as required by Section 65-40 of the Nurse 17 Practice Act; (v) the advanced practice registered nurse 18 19 must have completed the appropriate application 20 forms and paid the required fees as set by rule; 21 (vi) the advanced practice registered nurse 22 must provide evidence of satisfactory completion 23 least 45 graduate contact hours 24 pharmacology for any new license issued with 25 Schedule II authority after the effective date of

this amendatory Act of the 97th General Assembly;

and
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- 2 (vii) the advanced practice registered nurse 3 must annually complete 5 hours of continuing 4 education in pharmacology;
  - (2.5) with respect to advanced practice registered nurses certified as nurse practitioners, nurse midwives, or clinical nurse specialists who do not meet the requirements of Section 65-43 of the Nurse Practice Act practicing in a hospital affiliate,
    - (A) the advanced practice registered nurse certified as a nurse practitioner, nurse midwife, or clinical nurse specialist has been privileged to prescribe any Schedule II through V controlled substances by the hospital affiliate upon the recommendation of the appropriate physician committee of the hospital affiliate in accordance with Section 65-45 of the Nurse Practice Act, has completed the appropriate application forms, and has paid the required fees as set by rule; and
    - (B) an advanced practice registered nurse certified as a nurse practitioner, nurse midwife, or clinical nurse specialist has been privileged to prescribe any Schedule II controlled substances by the hospital affiliate upon the recommendation of the appropriate physician committee of the hospital affiliate, then the following conditions must be met:

Ţ	(1) specific schedule if controlled substances
2	by oral dosage or topical or transdermal
3	application may be designated, provided that the
4	designated Schedule II controlled substances are
5	routinely prescribed by advanced practice
6	registered nurses in their area of certification;
7	the privileging documents must identify the
8	specific Schedule II controlled substances by
9	either brand name or generic name; privileges to
10	prescribe or dispense Schedule II controlled
11	substances to be delivered by injection or other
12	route of administration may not be granted;
13	(ii) any privileges must be controlled
14	substances limited to the practice of the advanced
15	practice registered nurse;
16	(iii) any prescription must be limited to no
17	more than a 30-day supply;
18	(iv) the advanced practice registered nurse
19	must discuss the condition of any patients for
20	whom a controlled substance is prescribed monthly
21	with the appropriate physician committee of the
22	hospital affiliate or its physician designee; and
23	(v) the advanced practice registered nurse
24	must meet the education requirements of this
25	Section;

(3) with respect to animal euthanasia agencies, the

- euthanasia agency has obtained a license from the Department of Financial and Professional Regulation and obtained a registration number from the Department; or
  - (4) with respect to prescribing psychologists, the prescribing psychologist has been delegated authority to prescribe any nonnarcotic Schedule III through V controlled substances by a collaborating physician licensed to practice medicine in all its branches in accordance with Section 4.3 of the Clinical Psychologist Licensing Act, and the prescribing psychologist has completed the appropriate application forms and has paid the required fees as set by rule.
  - (b) The mid-level practitioner shall only be licensed to prescribe those schedules of controlled substances for which a licensed physician has delegated prescriptive authority, except that an animal euthanasia agency does not have any prescriptive authority and except that a physician assistant shall have prescriptive authority in accordance with the Physician Assistant Practice Act of 1987. An A physician assistant and an advanced practice registered nurse is are prohibited from prescribing medications and controlled substances not set forth in the required written delegation of authority or as authorized by their practice Act.
  - (c) Upon completion of all registration requirements, physician assistants, advanced practice registered nurses, and animal euthanasia agencies may be issued a mid-level

- 1 practitioner controlled substances license for Illinois.
- 2 (d) A collaborating physician may, but is not required to,
  3 delegate prescriptive authority to an advanced practice
  4 registered nurse as part of a written collaborative agreement,
  5 and the delegation of prescriptive authority shall conform to
- the requirements of Section 65-40 of the Nurse Practice Act.
- 7 (e) (Blank). A collaborating physician may, but is not
  8 required to, delegate prescriptive authority to a physician
  9 assistant as part of a written collaborative agreement, and
  10 the delegation of prescriptive authority shall conform to the
  11 requirements of Section 7.5 of the Physician Assistant
- 12 Practice Act of 1987.
- 13 (f) Nothing in this Section shall be construed to prohibit 14 generic substitution.
- 15 (Source: P.A. 99-173, eff. 7-29-15; 100-453, eff. 8-25-17;
- 16 100-513, eff. 1-1-18; 100-863, eff. 8-14-18.)

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                  Statutes amended in order of appearance
 3
      225 ILCS 95/4
                                  from Ch. 111, par. 4604
 4
      225 ILCS 95/5.5
 5
      225 ILCS 95/6
                                  from Ch. 111, par. 4606
 6
      225 ILCS 95/7
                                  from Ch. 111, par. 4607
      225 ILCS 95/7.5
 7
 8
      225 ILCS 95/7.7
 9
      225 ILCS 95/7.8 new
10
      225 ILCS 95/7.9 new
11
      225 ILCS 95/17
                                  from Ch. 111, par. 4617
      225 ILCS 95/21
                                  from Ch. 111, par. 4621
12
      225 ILCS 95/22.2
13
                                  from Ch. 111, par. 4622.2
      225 ILCS 95/22.3
14
                                  from Ch. 111, par. 4622.3
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      225 ILCS 95/22.6
                                  from Ch. 111, par. 4622.6
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      720 ILCS 570/303.05
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