

Registration and Regulation Committee

Adopted in House Comm. on May 19, 2004

	09300SB2253ham001 LRB093 15878 AMC 50122 a
1	AMENDMENT TO SENATE BILL 2253
2	AMENDMENT NO Amend Senate Bill 2253 by replacing
3	everything after the enacting clause with the following:
4	"Section 5. The Pharmacy Practice Act of 1987 is amended by
5	changing Section 3 as follows:
6	(225 ILCS 85/3) (from Ch. 111, par. 4123)
7	(Section scheduled to be repealed on January 1, 2008)
8	Sec. 3. Definitions. For the purpose of this Act, except
9	where otherwise limited therein:
10	(a) "Pharmacy" or "drugstore" means and includes every
11	store, shop, pharmacy department, or other place where
12	pharmaceutical care is provided by a pharmacist (1) where
13	drugs, medicines, or poisons are dispensed, sold or offered for
14	sale at retail, or displayed for sale at retail; or (2) where
15	prescriptions of physicians, dentists, veterinarians,
16	podiatrists, or therapeutically certified optometrists, within
17	the limits of their licenses, are compounded, filled, or
18	dispensed; or (3) which has upon it or displayed within it, or
19	affixed to or used in connection with it, a sign bearing the
20	word or words "Pharmacist", "Druggist", "Pharmacy",
21	"Pharmaceutical Care", "Apothecary", "Drugstore", "Medicine
22	Store", "Prescriptions", "Drugs", "Medicines", or any word or
23	words of similar or like import, either in the English language
24	or any other language; or (4) where the characteristic

prescription sign (Rx) or similar design is exhibited; or (5) any store, or shop, or other place with respect to which any of the above words, objects, signs or designs are used in any advertisement.

5 (b) "Drugs" means and includes (1) articles recognized in the official United States Pharmacopoeia/National Formulary 6 7 (USP/NF), or any supplement thereto and being intended for and 8 having for their main use the diagnosis, cure, mitigation, treatment or prevention of disease in man or other animals, as 9 10 approved by the United States Food and Drug Administration, but 11 does not include devices or their components, parts, or accessories; and (2) all other articles intended for and having 12 13 for their main use the diagnosis, cure, mitigation, treatment or prevention of disease in man or other animals, as approved 14 15 by the United States Food and Drug Administration, but does not 16 include devices or their components, parts, or accessories; and 17 (3) articles (other than food) having for their main use and 18 intended to affect the structure or any function of the body of man or other animals; and (4) articles having for their main 19 20 use and intended for use as a component or any articles 21 specified in clause (1), (2) or (3); but does not include 22 devices or their components, parts or accessories.

(c) "Medicines" means and includes all drugs intended for
human or veterinary use approved by the United States Food and
Drug Administration.

26 "Practice of pharmacy" means the provision (d) of 27 pharmaceutical care to patients as determined by the 28 pharmacist's professional judgment in the following areas, 29 which may include but are not limited to (1) patient 30 counseling, (2) interpretation and assisting in the monitoring 31 of appropriate drug use and prospective drug utilization 32 review, (3) providing information on the therapeutic values, 33 reactions, drug interactions, side effects, uses, selection of medications and medical devices, and outcome of drug therapy, 34

1 (4) participation in drug selection, drug monitoring, drug 2 utilization review, evaluation, administration, 3 interpretation, application of pharmacokinetic and laboratory 4 data to design safe and effective drug regimens, (5) drug 5 research (clinical and scientific), and (6) compounding and 6 dispensing of drugs and medical devices.

7 (e) "Prescription" means and includes any written, oral, 8 facsimile, or electronically transmitted order for drugs or medical devices, issued by a physician licensed to practice 9 medicine in all its branches, dentist, veterinarian, or 10 podiatrist, or therapeutically certified optometrist, within 11 the limits of their licenses, by a physician assistant in 12 accordance with subsection (f) of Section 4, or by an advanced 13 14 practice nurse in accordance with subsection (g) of Section 4, 15 containing the following: (1) name of the patient; (2) date when prescription was issued; (3) name and strength of drug or 16 17 description of the medical device prescribed; and (4) quantity, 18 (5) directions for use, (6) prescriber's name, address and 19 signature, and (7) DEA number where required, for controlled 20 substances. DEA numbers shall not be required on inpatient drug 21 orders.

(f) "Person" means and includes a natural person, copartnership, association, corporation, government entity, or any other legal entity.

25 (g) "Department" means the Department of Professional 26 Regulation.

(h) "Board of Pharmacy" or "Board" means the State Board ofPharmacy of the Department of Professional Regulation.

29 (i) "Director" means the Director of Professional30 Regulation.

31 (j) "Drug product selection" means the interchange for a 32 prescribed pharmaceutical product in accordance with Section 33 25 of this Act and Section 3.14 of the Illinois Food, Drug and 34 Cosmetic Act.

(k) "Inpatient drug order" means an order issued by an 1 2 authorized prescriber for a resident or patient of a facility 3 licensed under the Nursing Home Care Act or the Hospital Licensing Act, or "An Act in relation to the founding and 4 5 operation of the University of Illinois Hospital and the conduct of University of Illinois health care programs", 6 7 approved July 3, 1931, as amended, or a facility which is operated by the Department of Human Services (as successor to 8 of Mental Health 9 the Department and Developmental 10 Disabilities) or the Department of Corrections.

11 (k-5) "Pharmacist" means an individual health care 12 professional and provider currently licensed by this State to 13 engage in the practice of pharmacy.

(1) "Pharmacist in charge" means the licensed pharmacist whose name appears on a pharmacy license and who is responsible for all aspects of the operation related to the practice of pharmacy.

18 (m) "Dispense" means the delivery of drugs and medical 19 devices, in accordance with applicable State and federal laws 20 and regulations, to the patient or the patient's representative 21 authorized to these products, receive including the 22 preparation, compounding, packaging, and labeling necessary for delivery, interpretation, computer entry, and verification 23 24 of medication orders and prescriptions, drug product 25 selection, and any recommending or advising concerning the 26 contents and therapeutic values and uses thereof. "Dispense" does not mean the physical delivery to a patient or a patient's 27 28 representative in a home or institution by a designee of a 29 pharmacist or by common carrier. "Dispense" also does not mean 30 the physical delivery of a drug or medical device to a patient 31 or patient's representative by a pharmacist's designee within a 32 pharmacy or drugstore while the pharmacist is on duty and the 33 pharmacy is open.

34

(n) "Mail-order pharmacy" means a pharmacy that is located

in a state of the United States, other than Illinois, that 1 delivers, dispenses or distributes, through the United States 2 3 Postal Service or other common carrier, to Illinois residents, 4 any substance which requires a prescription.

5 (\circ) "Compounding" means the preparation, mixing, assembling, packaging, or labeling of a drug or medical device: 6 7 (1) as the result of a practitioner's prescription drug order 8 or initiative that is dispensed pursuant to a prescription in the course of professional practice; or (2) for the purpose of, 9 or incident to, research, teaching, or chemical analysis; or 10 in anticipation of prescription drug orders based on 11 (3) routine, regularly observed prescribing patterns. 12

13 "Confidential information" means information, (p) maintained by the pharmacist in the patient's records, released 14 15 only (i) to the patient or, as the patient directs, to other 16 practitioners and other pharmacists or (ii) to any other person authorized by law to receive the information. 17

18 "Prospective drug review" or "drug utilization (q) 19 evaluation" means a screening for potential drug therapy 20 problems due to therapeutic duplication, drug-disease 21 contraindications, drug-drug interactions (including serious 22 interactions with nonprescription or over-the-counter drugs), 23 drug-food interactions, incorrect drug dosage or duration of 24 drug treatment, drug-allergy interactions, and clinical abuse 25 or misuse.

26 (r) "Patient counseling" means the communication between a 27 pharmacist or a student pharmacist under the direct supervision 28 of a pharmacist and a patient or the patient's representative 29 about the patient's medication or device for the purpose of optimizing proper use of prescription medications or devices. 30 31 The offer to counsel by the pharmacist or the pharmacist's 32 designee, and subsequent patient counseling by the pharmacist 33 student pharmacist, shall be made in a face-to-face or communication with the patient or patient's representative 34

unless, in the professional judgment of the pharmacist, a 1 2 is face-to-face communication deemed inappropriate or 3 unnecessary. In that instance, the offer to counsel or patient 4 counseling may be made in a written communication, by 5 telephone, or in a manner determined by the pharmacist to be 6 appropriate.

7 (s) "Patient profiles" or "patient drug therapy record" 8 means the obtaining, recording, and maintenance of patient 9 prescription information, including prescriptions for 10 controlled substances, and personal information.

(t) "Pharmaceutical care" includes, but is not limited to, the act of monitoring drug use and other patient care services intended to achieve outcomes that improve the patient's quality of life but shall not include the sale of over-the-counter drugs by a seller of goods and services who does not dispense prescription drugs.

"Medical device" means an instrument, apparatus, 17 (u) 18 implement, machine, contrivance, implant, in vitro reagent, or 19 other similar or related article, including any component part 20 or accessory, required under federal law to bear the label 21 "Caution: Federal law requires dispensing by or on the order of a physician". A seller of goods and services who, only for the 22 purpose of retail sales, compounds, sells, rents, or leases 23 24 medical devices shall not, by reasons thereof, be required to 25 be a licensed pharmacy.

(v) "Unique identifier" means an electronic signature,
handwritten signature or initials, thumb print, or other
acceptable individual biometric or electronic identification
process as approved by the Department.

30 (Source: P.A. 92-880, eff. 1-1-04; 93-571, eff. 8-20-03.)

31 Section 99. Effective date. This Act takes effect upon 32 becoming law.".