By: Representative Powell

To: Public Health and Human Services; Insurance

HOUSE BILL NO. 1220

AN ACT TO AMEND SECTION 73-21-73, MISSISSIPPI CODE OF 1972, TO REVISE THE DEFINITION OF THE TERM "PRACTICE OF PHARMACY" IN THE PHARMACY PRACTICE ACT TO INCLUDE THE PROVIDING OF PATIENT CARE SERVICES; TO CREATE NEW SECTION 73-21-131, MISSISSIPPI CODE OF 5 1972, TO AUTHORIZE PHARMACISTS TO PROVIDE APPROVED PATIENT CARE 6 SERVICES IN ACCORDANCE WITH RULES ADOPTED BY THE STATE BOARD OF 7 PHARMACY AND PURSUANT TO A PROTOCOL; TO CREATE NEW SECTION 83-41-221, MISSISSIPPI CODE OF 1972, TO PROVIDE THAT WHENEVER ANY 8 9 INSURANCE POLICY ISSUED IN THIS STATE PROVIDES FOR PAYMENT OR 10 REIMBURSEMENT FOR ANY SERVICE THAT IS WITHIN THE LAWFUL SCOPE OF 11 PRACTICE OF A DULY LICENSED PHARMACIST, THE INSURED OR OTHER 12 PERSON ENTITLED TO BENEFITS UNDER THAT POLICY MAY BE PAID OR 13 REIMBURSED FOR THAT SERVICE WHEN THE SERVICE IS PERFORMED BY A DULY LICENSED PHARMACIST; AND FOR RELATED PURPOSES. 14

- BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF MISSISSIPPI:
- SECTION 1. Section 73-21-73, Mississippi Code of 1972, is
- 17 amended as follows:
- 18 73-21-73. As used in this chapter, unless the context
- 19 requires otherwise:
- 20 (a) "Administer" means the direct application of a
- 21 prescription drug pursuant to a lawful order of a practitioner to
- 22 the body of a patient by injection, inhalation, ingestion or any
- 23 other means.

- (b) "Board of Pharmacy," "Pharmacy Board," "MSBP" or
- 25 "board" means the State Board of Pharmacy.
- 26 (c) "Compounding" means (i) the production,
- 27 preparation, propagation, conversion or processing of a sterile or
- 28 nonsterile drug or device either directly or indirectly by
- 29 extraction from substances of natural origin or independently by
- 30 means of chemical or biological synthesis or from bulk chemicals
- 31 or the preparation, mixing, measuring, assembling, packaging or
- 32 labeling of a drug or device as a result of a practitioner's
- 33 prescription drug order or initiative based on the
- 34 practitioner/patient/pharmacist relationship in the course of
- 35 professional practice, or (ii) for the purpose of, as an incident
- 36 to, research, teaching or chemical analysis and not for sale or
- 37 dispensing. Compounding also includes the preparation of drugs or
- 38 devices in anticipation of prescription drug orders based on
- 39 routine regularly observed prescribing patterns.
- 40 (d) "Continuing education unit" means ten (10) clock
- 41 hours of study or other such activity as may be approved by the
- 42 board, including, but not limited to, all programs which have been
- 43 approved by the American Council on Pharmaceutical Education.
- 44 (e) "Deliver" or "delivery" means the actual,
- 45 constructive or attempted transfer in any manner of a drug or
- 46 device from one person to another, whether or not for a
- 47 consideration, including, but not limited to, delivery by mailing
- 48 or shipping.

49 (f) "Device" means an instrument, apparatus, implem	49	implement
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- 50 machine, contrivance, implant, in vitro reagent or other similar
- or related article, including any component part or accessory
- 52 which is required under federal or state law to be prescribed by a
- 53 practitioner and dispensed by a pharmacist.
- 54 (q) "Dispense" or "dispensing" means the interpretation
- of a valid prescription of a practitioner by a pharmacist and the
- 56 subsequent preparation of the drug or device for administration to
- 57 or use by a patient or other individual entitled to receive the
- 58 drug.
- 59 (h) "Distribute" means the delivery of a drug or device
- 60 other than by administering or dispensing to persons other than
- 61 the ultimate consumer.
- (i) "Drug" means:
- (i) Articles recognized as drugs in the official
- 64 United States Pharmacopeia, official National Formulary, official
- 65 Homeopathic Pharmacopeia, other drug compendium or any supplement
- 66 to any of them;
- 67 (ii) Articles intended for use in the diagnosis,
- 68 cure, mitigation, treatment or prevention of disease in man or
- 69 other animals;
- 70 (iii) Articles other than food intended to affect
- 71 the structure or any function of the body of man or other animals;
- 72 and

- 73 (iv) Articles intended for use as a component of
- 74 any articles specified in subparagraph (i), (ii) or (iii) of this
- 75 paragraph.
- 76 (j) "Drugroom" means a business, which does not require
- 77 the services of a pharmacist, where prescription drugs or
- 78 prescription devices are bought, sold, maintained or provided to
- 79 consumers.
- 80 (k) "Extern" means a student in the professional
- 81 program of a school of pharmacy accredited by the American Council
- 82 on Pharmaceutical Education who is making normal progress toward
- 83 completion of a professional degree in pharmacy.
- (1) "Foreign pharmacy graduate" means a person whose
- 85 undergraduate pharmacy degree was conferred by a recognized school
- 86 of pharmacy outside of the United States, the District of Columbia
- 87 and Puerto Rico. Recognized schools of pharmacy are those
- 88 colleges and universities listed in the World Health
- 89 Organization's World Directory of Schools of Pharmacy, or
- 90 otherwise approved by the Foreign Pharmacy Graduate Examination
- 91 Committee (FPGEC) certification program as established by the
- 92 National Association of Boards of Pharmacy.
- 93 (m) "Generic equivalent drug product" means a drug
- 94 product which (i) contains the identical active chemical
- 95 ingredient of the same strength, quantity and dosage form; (ii) is
- 96 of the same generic drug name as determined by the United States
- 97 Adoptive Names and accepted by the United States Food and Drug

- 98 Administration; and (iii) conforms to such rules and regulations
- 99 as may be adopted by the board for the protection of the public to
- 100 assure that such drug product is therapeutically equivalent.
- 101 (n) "Internet" means collectively the myriad of
- 102 computer and telecommunications facilities, including equipment
- 103 and operating software, which comprise the interconnected
- 104 worldwide network of networks that employ the Transmission Control
- 105 Protocol/Internet Protocol, or any predecessor or successor
- 106 protocol to such protocol, to communicate information of all kinds
- 107 by wire or radio.
- 108 (o) "Interested directly" means being employed by,
- 109 having full or partial ownership of, or control of, any facility
- 110 permitted or licensed by the Mississippi State Board of Pharmacy.
- 111 (p) "Interested indirectly" means having a spouse who
- is employed by any facility permitted or licensed by the
- 113 Mississippi State Board of Pharmacy.
- 114 (q) "Intern" means a person who has graduated from a
- 115 school of pharmacy but has not yet become licensed as a
- 116 pharmacist.
- 117 (r) "Manufacturer" means a person, business or other
- 118 entity engaged in the production, preparation, propagation,
- 119 conversion or processing of a prescription drug or device, if such
- 120 actions are associated with promotion and marketing of such drugs
- 121 or devices.

122	(s) "Manufacturer's distributor" means any person or
123	business who is not an employee of a manufacturer, but who
124	distributes sample drugs or devices, as defined under subsection
125	(i) of this section, under contract or business arrangement for a

126 manufacturer to practitioners.

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- (t) "Manufacturing" of prescription products means the production, preparation, propagation, conversion or processing of a drug or device, either directly or indirectly, by extraction from substances from natural origin or independently by means of chemical or biological synthesis, or from bulk chemicals and includes any packaging or repackaging of the substance(s) or labeling or relabeling of its container, if such actions are associated with promotion and marketing of such drug or devices.
- 135 (u) "Misappropriation of a prescription drug" means to
 136 illegally or unlawfully convert a drug, as defined in subsection
 137 (i) of this section, to one's own use or to the use of another.
- 138 (v) "Nonprescription drugs" means nonnarcotic medicines 139 or drugs that may be sold without a prescription and are 140 prepackaged and labeled for use by the consumer in accordance with 141 the requirements of the statutes and regulations of this state and 142 the federal government.
- 143 (w) "Person" means an individual, corporation, 144 partnership, association or any other legal entity.
- 145 (x) "Pharmacist" means an individual health care
 146 provider licensed by this state to engage in the practice of

- pharmacy. This recognizes a pharmacist as a learned professional who is authorized to provide patient services.
- 149 (y) "Pharmacy" means any location for which a pharmacy
 150 permit is required and in which prescription drugs are maintained,
 151 compounded and dispensed for patients by a pharmacist. This
 152 definition includes any location where pharmacy-related services
 153 are provided by a pharmacist.
- 154 (z) "Prepackaging" means the act of placing small
 155 precounted quantities of drug products in containers suitable for
 156 dispensing or administering in anticipation of prescriptions or
 157 orders.
- 158 (aa) "Unlawful or unauthorized possession" means
 159 physical holding or control by a pharmacist of a controlled
 160 substance outside the usual and lawful course of employment.
 - (bb) "Practice of pharmacy" means a health care service that includes, but is not limited to, the compounding, dispensing, and labeling of drugs or devices; interpreting and evaluating prescriptions; administering and distributing drugs and devices; the compounding, dispensing and labeling of drugs and devices; maintaining prescription drug records; advising and consulting concerning therapeutic values, content, hazards and uses of drugs and devices; initiating or modifying of drug therapy in accordance with written guidelines or protocols previously established and approved by the board; selecting drugs; participating in drug utilization reviews; storing prescription drugs and devices;

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- 172 ordering lab work in accordance with written guidelines or
- 173 protocols as defined by paragraph (11) of this section; providing
- 174 pharmacotherapeutic consultations; providing patient care services
- 175 as authorized under Section 73-21-131; supervising supportive
- 176 personnel; and such other acts, services, operations or
- 177 transactions necessary or incidental to the conduct of the
- 178 foregoing.
- 179 (cc) "Practitioner" means a physician, dentist,
- 180 veterinarian, or other health care provider authorized by law to
- 181 diagnose and prescribe drugs.
- 182 (dd) "Prescription" means a written, verbal or
- 183 electronically transmitted order issued by a practitioner for a
- 184 drug or device to be dispensed for a patient by a pharmacist.
- 185 (ee) "Prescription drug" or "legend drug" means a drug
- 186 which is required under federal law to be labeled with either of
- 187 the following statements prior to being dispensed or delivered:
- 188 (i) "Caution: Federal law prohibits dispensing
- 189 without prescription," or
- 190 (ii) "Caution: Federal law restricts this drug to
- 191 use by or on the order of a licensed veterinarian"; or a drug
- 192 which is required by any applicable federal or state law or
- 193 regulation to be dispensed on prescription only or is restricted
- 194 to use by practitioners only.

195		(ff)	"Product	selecti	ion"	means	the	disper	sing	of	a
196	generic	equivale	ent drug	product	in	lieu o	f the	drug	produ	ıct	
197	ordered	by the	orescribe	er.							

- 198 (gg) "Provider" or "primary health care provider"

 199 includes a pharmacist who provides health care services within his

 200 or her scope of practice pursuant to state law and regulation.
- (hh) "Registrant" means a pharmacy or other entity
 which is registered with the Mississippi State Board of Pharmacy
 to buy, sell or maintain controlled substances.
- (ii) "Repackager" means a person registered by the
 Federal Food and Drug Administration as a repackager who removes a
 prescription drug product from its marketed container and places
 it into another, usually of smaller size, to be distributed to
 persons other than the consumer.
- (jj) "Reverse distributor" means a business operator
 that is responsible for the receipt and appropriate return or
 disposal of unwanted, unneeded or outdated stocks of controlled or
 uncontrolled drugs from a pharmacy.
- (kk) "Supportive personnel" or "pharmacist technician"
 means those individuals utilized in pharmacies whose
 responsibilities are to provide nonjudgmental technical services
 concerned with the preparation and distribution of drugs under the
 direct supervision and responsibility of a pharmacist.
- 218 (11) "Written guideline or protocol" means an agreement 219 in which any practitioner authorized to prescribe drugs delegates

- 220 to a pharmacist authority to conduct specific prescribing
- 221 functions in an institutional setting, or with individual
- 222 patients, provided that a specific protocol agreement is signed on
- 223 each patient and is filed as required by law or by rule or
- 224 regulation of the board.
- 225 (mm) "Wholesaler" means a person who buys or otherwise
- 226 acquires prescription drugs or prescription devices for resale or
- 227 distribution, or for repackaging for resale or distribution, to
- 228 persons other than consumers.
- (nn) "Pharmacy benefit manager" has the same meaning as
- 230 defined in Section 73-21-153.
- 231 **SECTION 2.** The following shall be codified as Section
- 232 73-21-131, Mississippi Code of 1972:
- 233 73-21-131. In accordance with rules adopted by the State
- 234 Board of Pharmacy and pursuant to a protocol, a pharmacist may
- 235 provide approved patient care services including, but not limited
- 236 to, smoking cessation and travel health services.
- 237 **SECTION 3.** The following shall be codified as Section
- 238 83-41-221, Mississippi Code of 1972:
- 239 83-41-221. Whenever any policy of insurance or any medical
- 240 service plan or hospital service contract or hospital and medical
- 241 service contract issued in this state provides for payment or
- 242 reimbursement for any service that is within the lawful scope of
- 243 practice of a duly licensed pharmacist as defined in Section
- 244 73-21-73, the insured or other person entitled to benefits under

245	t.hat.	policy.	plan	or	contract	mav	be	paid	or	reimbursed	for	that

- 246 service when the service is performed by a duly licensed
- 247 pharmacist.
- 248 **SECTION 4.** This act shall take effect and be in force from
- 249 and after July 1, 2017.

