

By: Representatives Mims, Dixon, Currie

To: Public Health and Human Services

HOUSE BILL NO. 462

1 AN ACT TO AMEND SECTION 73-21-69, MISSISSIPPI CODE OF 1972,
2 TO EXTEND THE DATE OF THE REPEALER ON THE MISSISSIPPI PHARMACY
3 PRACTICE ACT AND INCLUDE ADDITIONAL SECTIONS IN THE REPEALER; TO
4 REENACT SECTIONS 73-21-71 THROUGH 73-21-123, MISSISSIPPI CODE OF
5 1972, WHICH ARE THE MISSISSIPPI PHARMACY PRACTICE ACT; TO AMEND
6 REENACTED SECTION 73-21-73, MISSISSIPPI CODE OF 1972, TO MAKE SOME
7 MINOR NONSUBSTANTIVE CHANGES; TO AMEND REENACTED SECTION 73-21-75,
8 MISSISSIPPI CODE OF 1972, TO PROVIDE THAT WHEN AN ELECTION IS
9 REQUIRED BY THE MISSISSIPPI PHARMACISTS ASSOCIATION TO NARROW THE
10 NUMBER OF POTENTIAL CANDIDATES FOR NOMINATIONS TO THE BOARD, THE
11 ASSOCIATION SHALL PROVIDE A BALLOT TO EACH LICENSED PHARMACIST; TO
12 AMEND REENACTED SECTION 73-21-83, MISSISSIPPI CODE OF 1972, WHICH
13 PROVIDES THAT THE STATE BOARD OF PHARMACY SHALL REGULATE THE
14 PRACTICE OF PHARMACY AND PHARMACY BENEFIT MANAGERS, TO DELETE THE
15 INDIVIDUAL REPEALER ON THAT SECTION; TO AMEND REENACTED SECTION
16 73-21-91, MISSISSIPPI CODE OF 1972, WHICH PROVIDES FOR LICENSE
17 RENEWAL FEES FOR PHARMACISTS AND PHARMACY BENEFIT MANAGERS, TO
18 DELETE THE INDIVIDUAL REPEALER ON THAT SECTION; TO AMEND REENACTED
19 SECTION 73-21-103, MISSISSIPPI CODE OF 1972, TO PROVIDE THAT THE
20 BOARD MAY IMPOSE A MONETARY PENALTY FOR ANY PERSON WHO USES THE
21 PRESCRIPTION MONITORING PROGRAM IN ANY MANNER OTHER THAN THAT FOR
22 WHICH IT WAS INTENDED; TO AMEND REENACTED SECTION 73-21-105,
23 MISSISSIPPI CODE OF 1972, TO AUTHORIZE THE BOARD TO DETERMINE THE
24 REGISTRATION PERIOD FOR FACILITIES THAT ENGAGE IN THE WHOLESALE
25 DISTRIBUTION OF PRESCRIPTION DRUGS AND REVERSE DISTRIBUTORS; TO
26 AMEND REENACTED SECTION 73-21-106, MISSISSIPPI CODE OF 1972, TO
27 PROVIDE THAT THE BOARD SHALL ESTABLISH THE CRITERIA THAT EACH
28 NONRESIDENT PHARMACY MUST MEET TO QUALIFY FOR A NONRESIDENT
29 PERMIT; TO PROVIDE THAT AFTER A PERMIT HAS BEEN ISSUED, IT MAY NOT
30 BE AMENDED, TRANSFERRED OR REASSIGNED; TO PROVIDE THAT A
31 PHARMACIST-IN-CHARGE OF A NONRESIDENT PHARMACY MAY NOT BE THE
32 PHARMACIST-IN-CHARGE AT ANY OTHER LOCATION THAT HAS BEEN ISSUED A
33 PERMIT BY THE BOARD; TO BRING FORWARD SECTION 73-21-125,
34 MISSISSIPPI CODE OF 1972, WHICH PROVIDES FOR CIVIL IMMUNITY FOR



35 COMMUNITY PHARMACIES AND PHARMACISTS WORKING IN THOSE PHARMACIES,
36 TO INCLUDE THE SECTION IN THE GENERAL REPEALER ON THE PHARMACY
37 PRACTICE ACT; TO AMEND SECTION 73-21-126, MISSISSIPPI CODE OF
38 1972, WHICH PROVIDES FOR REGULATION OF WHOLESALE DISTRIBUTORS,
39 CHAIN PHARMACY WAREHOUSES AND RE-PACKAGERS, TO INCLUDE THE SECTION
40 IN THE GENERAL REPEALER ON THE PHARMACY PRACTICE ACT AND MAKE SOME
41 MINOR NONSUBSTANTIVE CHANGES; TO AMEND SECTION 73-21-127,
42 MISSISSIPPI CODE OF 1972, WHICH ESTABLISH A CONTROLLED SUBSTANCES
43 PRESCRIPTION MONITORING PROGRAM (PMP), TO INCLUDE THAT SECTION IN
44 THE GENERAL REPEALER ON THE PHARMACY PRACTICE ACT AND DELETE THE
45 INDIVIDUAL REPEALER ON THE SECTION; TO PROVIDE THAT PHARMACISTS
46 LICENSED BY THE MISSISSIPPI BOARD OF PHARMACY MUST BE REGISTERED
47 USERS OF THE PMP; TO PROVIDE THAT THE PMP THROUGH THE BOARD MAY
48 ESTABLISH THE COST OF ADMINISTRATION, MAINTENANCE, AND OPERATION
49 OF THE PROGRAM AND CHARGE AGENCIES A FEE BASED ON A FORMULA TO BE
50 DETERMINED BY THE BOARD, ASSESS CHARGES FOR INFORMATION AND/OR
51 STATISTICAL DATA PROVIDED TO AGENCIES, INSTITUTIONS AND
52 INDIVIDUALS; TO PROVIDE THAT ANY MISUSE OF THE PMP IS SUBJECT TO
53 PENALTIES; TO PROVIDE THAT THE BOARD AND THE PMP SHALL BE IMMUNE
54 FROM CIVIL LIABILITY ARISING FROM INACCURACY OF ANY OF THE
55 INFORMATION SUBMITTED TO THE PROGRAM; TO AMEND SECTION 73-21-129,
56 MISSISSIPPI CODE OF 1972, WHICH REQUIRES DRUG MANUFACTURERS TO
57 ALLOW PHARMACIES TO RETURN OUTDATED DRUGS FOR CREDIT OR REFUND, TO
58 INCLUDE THAT SECTION IN THE GENERAL REPEALER ON THE PHARMACY
59 PRACTICE ACT AND DELETE THE INDIVIDUAL REPEALER ON THE SECTION;
60 AND FOR RELATED PURPOSES.

61 BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF MISSISSIPPI:

62 **SECTION 1.** Section 73-21-69, Mississippi Code of 1972, is
63 amended as follows:

64 73-21-69. Sections 73-21-71 through 73-21- * * * 129, which
65 create the State Board of Pharmacy and prescribe its duties and
66 powers, shall stand repealed on July 1, * * * 2020.

67 **SECTION 2.** Section 73-21-71, Mississippi Code of 1972, is
68 reenacted as follows:

69 73-21-71. This chapter shall be known as the "Mississippi
70 Pharmacy Practice Act."

71 **SECTION 3.** Section 73-21-73, Mississippi Code of 1972, is
72 reenacted and amended as follows:



73 73-21-73. As used in this chapter, unless the context
74 requires otherwise:

75 (a) "Administer" means the direct application of a
76 prescription drug pursuant to a lawful order of a practitioner to
77 the body of a patient by injection, inhalation, ingestion or any
78 other means.

79 (b) "Board of Pharmacy," "Pharmacy Board," "MSBP" or
80 "board" means the State Board of Pharmacy.

81 (c) "Compounding" means (i) the production,
82 preparation, propagation, conversion or processing of a sterile or
83 nonsterile drug or device either directly or indirectly by
84 extraction from substances of natural origin or independently by
85 means of chemical or biological synthesis or from bulk chemicals
86 or the preparation, mixing, measuring, assembling, packaging or
87 labeling of a drug or device as a result of a practitioner's
88 prescription drug order or initiative based on the
89 practitioner/patient/pharmacist relationship in the course of
90 professional practice, or (ii) for the purpose of, as an incident
91 to, research, teaching or chemical analysis and not for sale or
92 dispensing. Compounding also includes the preparation of drugs or
93 devices in anticipation of prescription drug orders based on
94 routine regularly observed prescribing patterns.

95 (d) "Continuing education unit" means ten (10) clock
96 hours of study or other such activity as may be approved by the



97 board, including, but not limited to, all programs which have been
98 approved by the American Council on Pharmaceutical Education.

99 (e) "Deliver" or "delivery" means the actual,
100 constructive or attempted transfer in any manner of a drug or
101 device from one person to another, whether or not for a
102 consideration, including, but not limited to, delivery by mailing
103 or shipping.

104 (f) "Device" means an instrument, apparatus, implement,
105 machine, contrivance, implant, in vitro reagent or other similar
106 or related article, including any component part or accessory
107 which is required under federal or state law to be prescribed by a
108 practitioner and dispensed by a pharmacist.

109 (g) "Dispense" or "dispensing" means the interpretation
110 of a valid prescription of a practitioner by a pharmacist and the
111 subsequent preparation of the drug or device for administration to
112 or use by a patient or other individual entitled to receive the
113 drug.

114 (h) "Distribute" means the delivery of a drug or device
115 other than by administering or dispensing to persons other than
116 the ultimate consumer.

117 (i) "Drug" means:

118 (i) Articles recognized as drugs in the official
119 United States Pharmacopeia, official National Formulary, official
120 Homeopathic Pharmacopeia, other drug compendium or any supplement
121 to any of them;



122 (ii) Articles intended for use in the diagnosis,
123 cure, mitigation, treatment or prevention of disease in man or
124 other animals;

125 (iii) Articles other than food intended to affect
126 the structure or any function of the body of man or other animals;
127 and

128 (iv) Articles intended for use as a component of
129 any articles specified in subparagraph (i), (ii) or (iii) of this
130 paragraph.

131 (j) "Drugroom" means a business, which does not require
132 the services of a pharmacist, where prescription drugs or
133 prescription devices are bought, sold, maintained or provided to
134 consumers.

135 (k) "Extern" means a student in the professional
136 program of a school of pharmacy accredited by the American Council
137 on Pharmaceutical Education who is making normal progress toward
138 completion of a professional degree in pharmacy.

139 (l) "Foreign pharmacy graduate" means a person whose
140 undergraduate pharmacy degree was conferred by a recognized school
141 of pharmacy outside of the United States, the District of Columbia
142 and Puerto Rico. Recognized schools of pharmacy are those
143 colleges and universities listed in the World Health
144 Organization's World Directory of Schools of Pharmacy, or
145 otherwise approved by the Foreign Pharmacy Graduate Examination



146 Committee (FPGEC) certification program as established by the
147 National Association of Boards of Pharmacy.

148 (m) "Generic equivalent drug product" means a drug
149 product which (i) contains the identical active chemical
150 ingredient of the same strength, quantity and dosage form; (ii) is
151 of the same generic drug name as determined by the United States
152 Adoptive Names and accepted by the United States Food and Drug
153 Administration; and (iii) conforms to such rules and regulations
154 as may be adopted by the board for the protection of the public to
155 assure that such drug product is therapeutically equivalent.

156 (n) "Internet" means collectively the myriad of
157 computer and telecommunications facilities, including equipment
158 and operating software, which comprise the interconnected
159 worldwide network of networks that employ the Transmission Control
160 Protocol/Internet Protocol, or any predecessor or successor
161 protocol to such protocol, to communicate information of all kinds
162 by wire or radio.

163 (o) "Interested directly" means being employed by,
164 having full or partial ownership of, or control of, any facility
165 permitted or licensed by the Mississippi State Board of Pharmacy.

166 (p) "Interested indirectly" means having a spouse who
167 is employed by any facility permitted or licensed by the
168 Mississippi State Board of Pharmacy.



169 (q) "Intern" means a person who has graduated from a
170 school of pharmacy but has not yet become licensed as a
171 pharmacist.

172 (r) "Manufacturer" means a person, business or other
173 entity engaged in the production, preparation, propagation,
174 conversion or processing of a prescription drug or device, if such
175 actions are associated with promotion and marketing of such drugs
176 or devices.

177 (s) "Manufacturer's distributor" means any person or
178 business who is not an employee of a manufacturer, but who
179 distributes sample drugs or devices, as defined under subsection
180 (i) of this section, under contract or business arrangement for a
181 manufacturer to practitioners.

182 (t) "Manufacturing" of prescription products means the
183 production, preparation, propagation, conversion or processing of
184 a drug or device, either directly or indirectly, by extraction
185 from substances from natural origin or independently by means of
186 chemical or biological synthesis, or from bulk chemicals and
187 includes any packaging or repackaging of the substance(s) or
188 labeling or relabeling of its container, if such actions are
189 associated with promotion and marketing of such drug or devices.

190 (u) "Misappropriation of a prescription drug" means to
191 illegally or unlawfully convert a drug, as defined in subsection
192 (i) of this section, to one's own use or to the use of another.



193 (v) "Nonprescription drugs" means nonnarcotic medicines
194 or drugs that may be sold without a prescription and are
195 prepackaged and labeled for use by the consumer in accordance with
196 the requirements of the statutes and regulations of this state and
197 the federal government.

198 (w) "Person" means an individual, corporation,
199 partnership, association or any other legal entity.

200 (x) "Pharmacist" means an individual health care
201 provider licensed by this state to engage in the practice of
202 pharmacy. This recognizes a pharmacist as a learned professional
203 who is authorized to provide patient services.

204 (y) "Pharmacy" means any location for which a pharmacy
205 permit is required and in which prescription drugs are maintained,
206 compounded and dispensed for patients by a pharmacist. This
207 definition includes any location where pharmacy-related services
208 are provided by a pharmacist.

209 (z) "Prepackaging" means the act of placing small
210 precounted quantities of drug products in containers suitable for
211 dispensing or administering in anticipation of prescriptions or
212 orders.

213 (aa) "Unlawful or unauthorized * * * possession" means
214 physical holding or control by a pharmacist of a controlled
215 substance outside the usual and lawful course of employment.

216 (bb) "Practice of pharmacy" means a health care service
217 that includes, but is not limited to, the compounding, dispensing,



218 and labeling of drugs or devices; interpreting and evaluating
219 prescriptions; administering and distributing drugs and devices;
220 the compounding, dispensing and labeling of drugs and devices;
221 maintaining prescription drug records; advising and consulting
222 concerning therapeutic values, content, hazards and uses of drugs
223 and devices; initiating or modifying of drug therapy in accordance
224 with written guidelines or protocols previously established and
225 approved by the board; selecting drugs; participating in drug
226 utilization reviews; storing prescription drugs and devices;
227 ordering lab work in accordance with written guidelines or
228 protocols as defined by paragraph (11) of this section; providing
229 pharmacotherapeutic consultations; supervising supportive
230 personnel and such other acts, services, operations or
231 transactions necessary or incidental to the conduct of the
232 foregoing.

233 (cc) "Practitioner" means a physician, dentist,
234 veterinarian, or other health care provider authorized by law to
235 diagnose and prescribe drugs.

236 (dd) "Prescription" means a written, verbal or
237 electronically transmitted order issued by a practitioner for a
238 drug or device to be dispensed for a patient by a pharmacist.

239 (ee) "Prescription drug" or "legend drug" means a drug
240 which is required under federal law to be labeled with either of
241 the following statements prior to being dispensed or delivered:



242 (i) "Caution: Federal law prohibits dispensing
243 without prescription," or

244 (ii) "Caution: Federal law restricts this drug to
245 use by or on the order of a licensed veterinarian"; or a drug
246 which is required by any applicable federal or state law or
247 regulation to be dispensed on prescription only or is restricted
248 to use by practitioners only.

249 (ff) "Product selection" means the dispensing of a
250 generic equivalent drug product in lieu of the drug product
251 ordered by the prescriber.

252 (gg) "Provider" or "primary health care provider"
253 includes a pharmacist who provides health care services within his
254 or her scope of practice pursuant to state law and regulation.

255 (hh) "Registrant" means a pharmacy or other entity
256 which is registered with the Mississippi State Board of Pharmacy
257 to buy, sell or maintain controlled substances.

258 (ii) "Repackager" means a person registered by the
259 Federal Food and Drug Administration as a repackager who removes a
260 prescription drug product from its marketed container and places
261 it into another, usually of smaller size, to be distributed to
262 persons other than the consumer.

263 (jj) "Reverse distributor" means a business operator
264 that is responsible for the receipt and appropriate return or
265 disposal of unwanted, unneeded or outdated stocks of controlled or
266 uncontrolled drugs from a pharmacy.



267 (kk) "Supportive personnel" or "pharmacist technician"
268 means those individuals utilized in pharmacies whose
269 responsibilities are to provide nonjudgmental technical services
270 concerned with the preparation and distribution of drugs under the
271 direct supervision and responsibility of a pharmacist.

272 (ll) "Written guideline or protocol" means an agreement
273 in which any practitioner authorized to prescribe drugs delegates
274 to a pharmacist authority to conduct specific prescribing
275 functions in an institutional setting, or with individual
276 patients, provided that a specific protocol agreement is signed on
277 each patient and is filed as required by law or by rule or
278 regulation of the board.

279 (mm) "Wholesaler" means a person who buys or otherwise
280 acquires prescription drugs or prescription devices for resale or
281 distribution, or for repackaging for resale or distribution, to
282 persons other than consumers.

283 (nn) "Pharmacy benefit manager" has the same meaning as
284 defined in Section 73-21-153.

285 **SECTION 4.** Section 73-21-75, Mississippi Code of 1972, is
286 reenacted and amended as follows:

287 73-21-75. (1) The State Board of Pharmacy created by former
288 Section 73-21-9 is * * * continued and reconstituted as follows:
289 The board shall consist of seven (7) appointed members. At least
290 one (1) appointment shall be made from each congressional
291 district. Each appointed member of the board shall be appointed



292 by the Governor, with the advice and consent of the Senate, from a
293 list of five (5) names submitted by the Mississippi Pharmacists
294 Association, with input from the Magnolia Pharmaceutical Society,
295 the Mississippi Independent Pharmacies Association (MIPA),
296 Mississippi Society of Health-System Pharmacists (MSHP) and
297 Mississippi College of Clinical Pharmacy (MCCP) and other
298 pharmacist associations or societies. Of the members appointed,
299 one (1) shall, at the time of appointment, have had five (5)
300 years' experience as a pharmacist at a facility holding an
301 institutional permit, and one (1) shall, at the time of
302 appointment, have had five (5) years' experience as a pharmacist
303 at a facility holding a retail permit. Any person appointed to
304 the board shall be limited to two (2) full terms of office during
305 any fifteen-year period, including any member serving on May 14,
306 1992.

307 (2) The members of the board appointed and serving prior to
308 July 1, 1983, whose terms have not expired by July 1, 1983, shall
309 serve the balance of their terms as members of the reconstituted
310 board, and they shall be considered to be from the same
311 congressional districts from which they were originally appointed
312 if they still reside therein, even if the district boundaries have
313 changed subsequent to their original appointments. The Governor
314 shall appoint the remaining members of the reconstituted board in
315 the manner prescribed in subsection (1) of this section on July 1,



316 1983. The initial members of the reconstituted board shall serve
317 terms of office as follows:

318 (a) The term of the member from the First Congressional
319 District shall expire on July 1, 1984; and from and after July 1,
320 1996, this appointment shall be designated as Post 1.

321 (b) The term of the member from the Second
322 Congressional District shall expire on July 1, 1988; and from and
323 after July 1, 1996, this appointment shall be designated as Post
324 2.

325 (c) The term of the member from the Third Congressional
326 District shall expire on July 1, 1986; and from and after July 1,
327 1996, this appointment shall be designated as Post 3.

328 (d) The term of the member from the Fourth
329 Congressional District shall expire on July 1, 1985; and from and
330 after July 1, 1996, this appointment shall be designated as Post
331 4.

332 (e) The term of the member from the Fifth Congressional
333 District shall expire on July 1, 1987; and from and after July 1,
334 1996, this appointment shall be designated as Post 5.

335 (f) The term of one (1) of the members from the state
336 at large shall expire on July 1, 1985; and from and after July 1,
337 1996, this appointment shall be designated as Post 6.

338 (g) The term of the other member from the state at
339 large shall expire on July 1, 1988; and from and after July 1,
340 1996, this appointment shall be designated as Post 7.



341 The appointments of members from congressional districts as
342 provided under this section shall be made from the congressional
343 districts as they existed on July 1, 2001.

344 (3) At the expiration of a term, members of the board shall
345 be appointed in the manner prescribed in subsection (1) of this
346 section for terms of five (5) years from the expiration date of
347 the previous terms. Any vacancy on the board prior to the
348 expiration of a term for any reason, including resignation,
349 removal, disqualification, death or disability, shall be filled by
350 appointment of the Governor in the manner prescribed in subsection
351 (1) of this section for the balance of the unexpired term. The
352 Mississippi Pharmacists Association, with input from the Magnolia
353 Pharmaceutical Society, the Mississippi Independent Pharmacies
354 Association (MIPA), Mississippi Society of Health-System
355 Pharmacists (MSHP) and Mississippi College of Clinical Pharmacy
356 (MCCP) and other pharmacist associations or societies, shall
357 submit a list of nominees no more than thirty (30) days after a
358 vacancy occurs, and the Governor shall fill such vacancies within
359 ninety (90) days after each such vacancy occurs. If an election
360 is required to narrow the number of potential candidates for
361 nominations to the board, the Mississippi Pharmacists Association
362 shall provide a ballot to each pharmacist holding a valid
363 Mississippi license.

364 (4) To be qualified to be a member of the board, a person
365 shall:



366 (a) Be an adult citizen of Mississippi for a period of
367 at least five (5) years preceding his appointment to the board;

368 (b) Be a pharmacist licensed and in good standing to
369 practice pharmacy in the State of Mississippi; and

370 (c) Have actively engaged in the practice of pharmacy
371 in Mississippi for a period of at least five (5) years.

372 (5) The Governor may remove any or all members of the board
373 on proof of unprofessional conduct, continued absence from the
374 state, or for failure to perform the duties of his office. Any
375 member who shall not attend two (2) consecutive meetings of the
376 board for any reason other than illness of such member shall be
377 subject to removal by the Governor. The president of the board
378 shall notify the Governor in writing when any such member has
379 failed to attend two (2) consecutive regular meetings. No removal
380 shall be made without first giving the accused an opportunity to
381 be heard in refutation of the charges made against him, and he
382 shall be entitled to receive a copy of the charges at the time of
383 filing.

384 **SECTION 5.** Section 73-21-77, Mississippi Code of 1972, is
385 reenacted as follows:

386 73-21-77. (1) Each person appointed as a member of the
387 board shall qualify by taking the oath prescribed by the
388 Constitution for the state officers, and shall file certificate
389 thereof in the Office of the Secretary of State within fifteen
390 (15) days after his appointment.



391 (2) There shall be a president of the board and such other
392 officers as deemed necessary by the board elected by and from its
393 membership.

394 (3) The board shall meet at least once each quarter to
395 transact business, and may meet at such additional times as it may
396 deem necessary. Such additional meetings may be called by the
397 president of the board or a majority of the members of the board.

398 (4) The place for each meeting shall be determined prior to
399 giving notice of such meeting and shall not be changed after such
400 notice is given without adequate subsequent notice.

401 (5) A majority of the members of the board shall constitute
402 a quorum for the conduct of the meeting and all actions of the
403 board shall be by a majority.

404 (6) Each member of the board shall receive a per diem as
405 provided in Section 25-3-69, not to exceed thirty (30) days in any
406 one (1) period of twelve (12) months, for each day actually
407 engaged in meetings of the board, together with necessary
408 traveling and other expenses as provided in Section 25-3-41.

409 **SECTION 6.** Section 73-21-79, Mississippi Code of 1972, is
410 reenacted as follows:

411 73-21-79. (1) The board shall employ an executive director
412 of the board. The executive director shall be a citizen of
413 Mississippi and a pharmacist licensed and in good standing to
414 practice pharmacy in the State of Mississippi, who has had five
415 (5) years' experience as a pharmacist.



416 (2) The executive director shall receive a salary to be set
417 by the board, subject to the approval of the State Personnel
418 Board, and shall be entitled to necessary expenses incurred in the
419 performance of his official duties. He shall devote full time to
420 the duties of his office and shall not be engaged in any other
421 business that will interfere with the duties of his office.

422 (3) The duties and responsibilities of the executive
423 director shall be defined by rules and regulations prescribed by
424 the board.

425 (4) The board may, in its discretion, employ persons in
426 addition to the executive director in such other positions or
427 capacities as it deems necessary to the proper conduct of board
428 business. Any pharmacist-investigator employed by the board may
429 have other part-time employment, provided that he shall not accept
430 any employment that would cause a conflict of interest in his
431 pharmacist-investigator duties. The board may employ legal
432 counsel to assist in the conduct of its business.

433 **SECTION 7.** Section 73-21-81, Mississippi Code of 1972, is
434 reenacted as follows:

435 73-21-81. The responsibility for the enforcement of the
436 provisions of this chapter shall be vested in the board. The
437 board shall have all of the duties, powers and authority
438 specifically granted by and necessary to the enforcement of this
439 chapter. The board may make, adopt, amend and repeal such rules
440 and regulations as may be deemed necessary by the board from time



441 to time for the proper administration and enforcement of this
442 chapter, in accordance with the provisions of the Mississippi
443 Administrative Procedures Law (Section 25-43-1 et seq.).

444 **SECTION 8.** Section 73-21-83, Mississippi Code of 1972, is
445 reenacted and amended as follows:

446 73-21-83. (1) The board shall be responsible for the
447 control and regulation of the practice of pharmacy, to include the
448 regulation of pharmacy externs or interns and pharmacist
449 technicians, in this state, the regulation of the wholesaler
450 distribution of drugs and devices as defined in Section 73-21-73,
451 the distribution of sample drugs or devices by manufacturer's
452 distributors as defined in Section 73-21-73 by persons other than
453 the original manufacturer or distributor in this state and the
454 regulation of pharmacy benefit managers as defined in Section
455 73-21-153.

456 (2) A license for the practice of pharmacy shall be obtained
457 by all persons prior to their engaging in the practice of
458 pharmacy. However, the provisions of this chapter shall not apply
459 to physicians, dentists, veterinarians, osteopaths or other
460 practitioners of the healing arts who are licensed under the laws
461 of the State of Mississippi and are authorized to dispense and
462 administer prescription drugs in the course of their professional
463 practice.

464 (3) The initial licensure fee shall be set by the board but
465 shall not exceed Two Hundred Dollars (\$200.00), except the initial



466 licensure fee for pharmacy benefit managers shall be set by the
467 board but shall not exceed Five Hundred Dollars (\$500.00).

468 (4) All students actively enrolled in a professional school
469 of pharmacy accredited by the American Council on Pharmaceutical
470 Education who are making satisfactory progress toward graduation
471 and who act as an extern or intern under the direct supervision of
472 a pharmacist in a location permitted by the Board of Pharmacy must
473 obtain a pharmacy student registration prior to engaging in such
474 activity. The student registration fee shall be set by the board
475 but shall not exceed One Hundred Dollars (\$100.00).

476 (5) All persons licensed to practice pharmacy prior to July
477 1, 1991, by the State Board of Pharmacy under Section 73-21-89
478 shall continue to be licensed under the provisions of Section
479 73-21-91.

480 * * *

481 **SECTION 9.** Section 73-21-85, Mississippi Code of 1972, is
482 reenacted as follows:

483 73-21-85. (1) To obtain a license to engage in the practice
484 of pharmacy by examination, or by score transfer, the applicant
485 shall:

486 (a) Have submitted a written application on the form
487 prescribed by the board;

488 (b) Be of good moral character;



489 (c) Have graduated from a school or college of pharmacy
490 accredited by the American Council of Pharmaceutical Education and
491 have been granted a pharmacy degree therefrom;

492 (d) Have successfully passed an examination approved by
493 the board;

494 (e) Have paid all fees specified by the board for
495 examination, not to exceed the cost to the board of administering
496 the examination;

497 (f) Have paid all fees specified by the board for
498 licensure; and

499 (g) Have submitted evidence of externship and/or
500 internship as specified by the board.

501 (2) To obtain a license to engage in the practice of
502 pharmacy, a foreign pharmacy graduate applicant shall obtain the
503 National Association of Boards of Pharmacy's Foreign Pharmacy
504 Graduate Examination Committee's certification, which shall
505 include, but not be limited to, successfully passing the Foreign
506 Pharmacy Graduate Equivalency Examination and attaining a total
507 score of at least five hundred fifty (550) on the Test of English
508 as a Foreign Language (TOEFL), and shall:

509 (a) Have submitted a written application on the form
510 prescribed by the board;

511 (b) Be of good moral character;

512 (c) Have graduated and been granted a pharmacy degree
513 from a college or school of pharmacy recognized and approved by



514 the National Association of Boards of Pharmacy's Foreign Pharmacy
515 Graduate Examination Committee;

516 (d) Have paid all fees specified by the board for
517 examination, not to exceed the cost to the board of administering
518 the examination;

519 (e) Have successfully passed an examination approved by
520 the board;

521 (f) Have completed the number of internship hours as
522 set forth by regulations of the board; and

523 (g) Have paid all fees specified by the board for
524 licensure.

525 (3) Each application or filing made under this section shall
526 include the social security number(s) of the applicant in
527 accordance with Section 93-11-64.

528 (4) To insure that all applicants are of good moral
529 character, the board shall conduct a criminal history records
530 check on all applicants for a license. In order to determine the
531 applicant's suitability for licensing, the applicant shall be
532 fingerprinted. The board shall submit the fingerprints to the
533 Department of Public Safety for a check of the state criminal
534 records and forwarded to the Federal Bureau of Investigation for a
535 check of the national criminal records. The Department of Public
536 Safety shall disseminate the results of the state check and the
537 national check to the board for a suitability determination. The
538 board shall be authorized to collect from the applicant the amount



539 of the fee that the Department of Public Safety charges the board
540 for the fingerprinting, whether manual or electronic, and the
541 state and national criminal history records checks.

542 (5) To insure that all applicants are of good moral
543 character, the board, upon request of the Dean of the University
544 of Mississippi School of Pharmacy, shall be authorized to conduct
545 a criminal history records check on all applicants for enrollment
546 into the School of Pharmacy. In order to determine the
547 applicant's suitability for enrollment and licensing, the
548 applicant shall be fingerprinted. The board shall submit the
549 fingerprints to the Department of Public Safety for a check of the
550 state criminal records and forwarded to the Federal Bureau of
551 Investigation for a check of the national criminal records. The
552 Department of Public Safety shall disseminate the results of the
553 state check and the national check to the board for a suitability
554 determination and the board shall forward the results to the Dean
555 of the School of Pharmacy. The board shall be authorized to
556 collect from the applicant the amount of the fee that the
557 Department of Public Safety charges the board for the
558 fingerprinting, whether manual or electronic, and the state and
559 national criminal history records checks.

560 **SECTION 10.** Section 73-21-87, Mississippi Code of 1972, is
561 reenacted as follows:



562 73-21-87. (1) To obtain a license to engage in the practice
563 of pharmacy by reciprocity or license transfer, the applicant
564 shall:

565 (a) Have submitted a written application on the form
566 prescribed by the board;

567 (b) Be of good moral character;

568 (c) Have possessed at the time of initial licensure as
569 a pharmacist such other qualifications necessary to have been
570 eligible for licensure at that time in that state;

571 (d) Have presented to the board proof that any license
572 or licenses granted to the applicant by any other states have not
573 been suspended, revoked, cancelled or otherwise restricted for any
574 reason except nonrenewal or the failure to obtain required
575 continuing education credits; and

576 (e) Have paid all fees specified by the board for
577 licensure.

578 (2) No applicant shall be eligible for licensure by
579 reciprocity or license transfer unless the state in which the
580 applicant was initially licensed also grants a reciprocal license
581 or transfer license to pharmacists licensed by this state under
582 like circumstances and conditions.

583 (3) The issuance of a license by reciprocity to a
584 military-trained applicant or military spouse shall be subject to
585 the provisions of Section 73-50-1.



586 (4) Each application or filing made under this section shall
587 include the social security number(s) of the applicant in
588 accordance with Section 93-11-64.

589 **SECTION 11.** Section 73-21-89, Mississippi Code of 1972, is
590 reenacted as follows:

591 73-21-89. (1) The board shall issue a license to practice
592 pharmacy to any person, if such person be otherwise qualified,
593 upon presentation to the board of:

594 (a) Satisfactory proof that the applicant has been
595 graduated from the University of Mississippi School of Pharmacy;

596 (b) Written application for licensure; and

597 (c) Payment of all fees specified by the board for
598 licensure.

599 (2) The board shall not issue any new licenses pursuant to
600 this section after June 30, 1987.

601 (3) Each application or filing made under this section shall
602 include the social security number(s) of the applicant in
603 accordance with Section 93-11-64, Mississippi Code of 1972.

604 **SECTION 12.** Section 73-21-91, Mississippi Code of 1972, is
605 reenacted and amended as follows:

606 73-21-91. (1) Every pharmacist shall renew his license
607 annually. To renew his license, a pharmacist shall:

608 (a) Submit an application for renewal on the form
609 prescribed by the board;



610 (b) Submit satisfactory evidence of the completion in
611 the last licensure period of such continuing education units as
612 shall be required by the board, but in no case less than one (1)
613 continuing education unit in the last licensure period;

614 (c) (i) Pay any renewal fees as required by the board,
615 not to exceed One Hundred Dollars (\$100.00) for each annual
616 licensing period, provided that the board may add a surcharge of
617 not more than Five Dollars (\$5.00) to a license renewal fee to
618 fund a program to aid impaired pharmacists or pharmacy students.
619 Any pharmacist license renewal received postmarked after December
620 31 of the renewal period will be returned and a Fifty Dollar
621 (\$50.00) late renewal fee will be assessed before renewal.

622 (ii) The license fee for a pharmacy benefit
623 manager shall be set by the board, but shall not exceed Five
624 Hundred Dollars (\$500.00). Any license renewal received
625 postmarked after December 31 of the renewal period will be
626 returned and a Five Hundred Dollar (\$500.00) late renewal fee will
627 be assessed before renewal.

628 (2) Any pharmacist who has defaulted in license renewal may
629 be reinstated within two (2) years upon payment of renewal fees in
630 arrears and presentation of evidence of the required continuing
631 education. Any pharmacist defaulting in license renewal for a
632 period in excess of two (2) years shall be required to
633 successfully complete the examination given by the board pursuant
634 to Section 73-21-85 before being eligible for reinstatement as a



635 pharmacist in Mississippi, or shall be required to appear before
636 the board to be examined for his competence and knowledge of the
637 practice of pharmacy, and may be required to submit evidence of
638 continuing education. If the person is found fit by the board to
639 practice pharmacy in this state, the board may reinstate his
640 license to practice pharmacy upon payment of all renewal fees in
641 arrears.

642 (3) Each application or filing made under this section shall
643 include the social security number(s) of the applicant in
644 accordance with Section 93-11-64.

645 * * *

646 **SECTION 13.** Section 73-21-93, Mississippi Code of 1972, is
647 reenacted as follows:

648 73-21-93. (1) The examination for licensure required under
649 Section 73-21-85 shall be given by the board at least once during
650 each year. The board shall determine the content and subject
651 matter of each examination, the place, time and date of the
652 administration of the examination and those persons who have
653 successfully passed the examination.

654 (2) The examination shall be prepared to measure the
655 competence of the applicant to engage in the practice of pharmacy.
656 The board may employ and cooperate with any organization or
657 consultant in the preparation and grading of an appropriate
658 examination, but shall retain the sole discretion and



659 responsibility of determining which applicants have successfully
660 passed such an examination.

661 (3) The board shall have authority to use the laboratories
662 of the school of pharmacy and other facilities of the University
663 of Mississippi for the purpose of examining applicants.

664 **SECTION 14.** Section 73-21-95, Mississippi Code of 1972, is
665 reenacted as follows:

666 73-21-95. The assistant pharmacist license is hereby
667 abolished after April 30, 1984. The board shall issue a license
668 to practice pharmacy to those persons presently holding an
669 assistant pharmacist license upon their meeting the requirements
670 of Section 73-21-91.

671 **SECTION 15.** Section 73-21-97, Mississippi Code of 1972, is
672 reenacted as follows:

673 73-21-97. (1) The board may refuse to issue or renew, or
674 may suspend, reprimand, revoke or restrict the license,
675 registration or permit of any person upon one or more of the
676 following grounds:

677 (a) Unprofessional conduct as defined by the rules and
678 regulations of the board;

679 (b) Incapacity of a nature that prevents a pharmacist
680 from engaging in the practice of pharmacy with reasonable skill,
681 confidence and safety to the public;

682 (c) Being found guilty by a court of competent
683 jurisdiction of one or more of the following:



684 (i) A felony;

685 (ii) Any act involving moral turpitude or gross
686 immorality; or

687 (iii) Violation of pharmacy or drug laws of this
688 state or rules or regulations pertaining thereto, or of statutes,
689 rules or regulations of any other state or the federal government;

690 (d) Fraud or intentional misrepresentation by a
691 licensee or permit holder in securing the issuance or renewal of a
692 license or permit;

693 (e) Engaging or aiding and abetting an individual to
694 engage in the practice of pharmacy without a license;

695 (f) Violation of any of the provisions of this chapter
696 or rules or regulations adopted pursuant to this chapter;

697 (g) Failure to comply with lawful orders of the board;

698 (h) Negligently or willfully acting in a manner
699 inconsistent with the health or safety of the public;

700 (i) Addiction to or dependence on alcohol or controlled
701 substances or the unauthorized use or possession of controlled
702 substances;

703 (j) Misappropriation of any prescription drug;

704 (k) Being found guilty by the licensing agency in
705 another state of violating the statutes, rules or regulations of
706 that jurisdiction;

707 (l) The unlawful or unauthorized possession of a
708 controlled substance;



709 (m) Willful failure to submit drug monitoring
710 information or willful submission of incorrect dispensing
711 information as required by the Prescription Monitoring Program
712 under Section 73-21-127;

713 (n) Failure to obtain the license, registration or
714 permit required by this chapter; or

715 (o) Violation(s) of the provisions of Sections 41-121-1
716 through 41-121-9 relating to deceptive advertisement by health
717 care practitioners. This paragraph shall stand repealed on July
718 1, 2016.

719 (2) In lieu of suspension, revocation or restriction of a
720 license as provided for above, the board may warn or reprimand the
721 offending pharmacist.

722 (3) In addition to the grounds specified in subsection (1)
723 of this section, the board shall be authorized to suspend the
724 license, registration or permit of any person for being out of
725 compliance with an order for support, as defined in Section
726 93-11-153. The procedure for suspension of a license,
727 registration or permit for being out of compliance with an order
728 for support, and the procedure for the reissuance or reinstatement
729 of a license, registration or permit suspended for that purpose,
730 and the payment of any fees for the reissuance or reinstatement of
731 a license, registration or permit suspended for that purpose,
732 shall be governed by Section 93-11-157 or 93-11-163, as the case
733 may be. If there is any conflict between any provision of Section



734 93-11-157 or 93-11-163 and any provision of this chapter, the
735 provisions of Section 93-11-157 or 93-11-163, as the case may be,
736 shall control.

737 **SECTION 16.** Section 73-21-99, Mississippi Code of 1972, is
738 reenacted as follows:

739 73-21-99. (1) Disciplinary action by the board against a
740 licensee, registrant or permit holder, or license, registration or
741 permit shall require the following:

742 (a) A sworn affidavit filed with the board charging a
743 licensee or permit holder with an act which is grounds for
744 disciplinary action as provided in Section 73-21-97; and

745 (b) An order of the Investigations Review Committee of
746 the board which shall cause the executive director of the board to
747 fix a time and place for a hearing by the board. The executive
748 director shall cause a written notice specifying the offense or
749 offenses for which the licensee or permit holder is charged and
750 notice of the time and place of the hearing to be served upon the
751 licensee or permit holder at least thirty (30) days prior to the
752 hearing date. Such notice may be served by mailing a copy thereof
753 by certified mail, postage prepaid, to the last-known residence or
754 business address of the licensee or permit holder.

755 (2) The board shall designate two (2) of its members to
756 serve on a rotating no longer than three-consecutive-month basis
757 with the executive director and legal counsel for the board as an
758 Investigations Review Committee, and the board's investigators



759 shall provide status reports solely to the Investigations Review
760 Committee during monthly meetings of the board. Such reports
761 shall be made on all on-going investigations, and shall apply to
762 any routine inspections which may give rise to the filing of a
763 complaint. In the event any complaint on a licensee comes before
764 the board for possible disciplinary action, the members of the
765 board serving on the Investigations Review Committee which
766 reviewed the investigation of such complaint shall recuse
767 themselves and not participate in the disciplinary proceeding.

768 (3) The board acting by and through its Investigation Review
769 Committee may, if deemed necessary, issue a letter of reprimand to
770 any licensee, registrant or permit holder in lieu of formal action
771 by the board.

772 (4) The board, acting by and through its executive director,
773 is hereby authorized and empowered to issue subpoenas for the
774 attendance of witnesses and the production of books and papers at
775 such hearing. Process issued by the board shall extend to all
776 parts of the state and shall be served by any person designated by
777 the board for such service.

778 (5) The accused shall have the right to appear either
779 personally or by counsel, or both, to produce witnesses or
780 evidence in his behalf, to cross-examine witnesses, and to have
781 subpoenas issued by the board.

782 (6) At the hearing, the board shall administer oaths as may
783 be necessary for the proper conduct of the hearing. All hearings



784 shall be conducted by the board, which shall not be bound by
785 strict rules of procedure or by the laws of evidence in the
786 conduct of its proceedings, but the determination shall be based
787 upon sufficient evidence to sustain it.

788 (7) Where, in any proceeding before the board, any witness
789 fails or refuses to attend upon a subpoena issued by the board,
790 refuses to testify, or refuses to produce any books and papers the
791 production of which is called for by a subpoena, the attendance of
792 such witness, the giving of his testimony or the production of the
793 books and papers shall be enforced by any court of competent
794 jurisdiction of this state in the manner provided for the
795 enforcement of attendance and testimony of witnesses in civil
796 cases in the courts of this state.

797 (8) The board shall, within thirty (30) days after
798 conclusion of the hearing, reduce its decision to writing and
799 forward an attested true copy thereof to the last-known residence
800 or business address of such licensee or permit holder by way of
801 United States first-class, certified mail, postage prepaid.

802 **SECTION 17.** Section 73-21-101, Mississippi Code of 1972, is
803 reenacted as follows:

804 73-21-101. (1) The right to appeal from the action of the
805 board in denying, revoking, suspending or refusing to renew any
806 license, registration or permit issued by the board, or fining or
807 otherwise disciplining any person is hereby granted. Such appeal
808 shall be to the chancery court of the county of the residence of



809 the licensee or permit holder on the record made, including a
810 verbatim transcript of the testimony at the hearing. The appeal
811 shall be taken within thirty (30) days after notice of the action
812 of the board in denying, revoking, suspending or refusing to renew
813 the license or permit, or fining or otherwise disciplining the
814 person. The appeal shall be perfected upon filing notice of the
815 appeal and by the prepayment of all costs, including the cost of
816 the preparation of the record of the proceedings by the board, and
817 the filing of a bond in the sum of Two Hundred Dollars (\$200.00),
818 conditioned that if the action of the board in denying, revoking,
819 suspending or refusing to renew the license or permit, or fining
820 or otherwise disciplining the person, be affirmed by the chancery
821 court, the licensee or permit holder will pay the costs of the
822 appeal and the action in the chancery court.

823 (2) If there is an appeal, such appeal shall act as a
824 supersedeas. The chancery court shall dispose of the appeal and
825 enter its decision promptly. The hearing on the appeal may, in
826 the discretion of the chancellor, be tried in vacation. The scope
827 of review of the chancery court shall be limited to a review of
828 the record made before the board to determine if the action of the
829 board is unlawful for the reason that it was (a) not supported by
830 substantial evidence, (b) arbitrary or capricious, (c) beyond the
831 power of the board to make, or (d) in violation of some statutory
832 or constitutional right of the appellant. The decision of the



833 chancery court may be appealed to the Supreme Court in the manner
834 provided by law.

835 (3) Actions taken by the board in suspending a license,
836 registration or permit when required by Section 93-11-157 or
837 93-11-163 are not actions from which an appeal may be taken under
838 this section. Any appeal of a suspension of a license,
839 registration or permit that is required by Section 93-11-157 or
840 93-11-163 shall be taken in accordance with the appeal procedure
841 specified in Section 93-11-157 or 93-11-163, as the case may be,
842 rather than the procedure specified in this section.

843 **SECTION 18.** Section 73-21-103, Mississippi Code of 1972, is
844 reenacted and amended as follows:

845 73-21-103. (1) Upon the finding of the existence of grounds
846 for action against any permitted facility or discipline of any
847 person holding a license, registration or permit, seeking a
848 license, registration or permit, seeking to renew a license or
849 permit under the provisions of this chapter, or practicing or
850 doing business without a license, registration or permit, the
851 board may impose one or more of the following penalties:

852 (a) Suspension of the offender's license, registration
853 and/or permit for a term to be determined by the board;

854 (b) Revocation of the offender's license, registration
855 and/or permit;

856 (c) Restriction of the offender's license, registration
857 and/or permit to prohibit the offender from performing certain



858 acts or from engaging in the practice of pharmacy in a particular
859 manner for a term to be determined by the board;

860 (d) Imposition of a monetary penalty as follows:

861 (i) For the first violation, a monetary penalty of
862 not less than Two Hundred Fifty Dollars (\$250.00) nor more than
863 One Thousand Dollars (\$1,000.00) for each violation;

864 (ii) For the second violation and subsequent
865 violations, a monetary penalty of not less than Five Hundred
866 Dollars (\$500.00) nor more than Five Thousand Dollars (\$5,000.00)
867 for each violation.

868 Money collected by the board under paragraph (d)(i), (ii) and
869 (iv) of this section shall be deposited to the credit of the State
870 General Fund of the State Treasury;

871 (iii) The board may assess a monetary penalty for
872 those reasonable costs that are expended by the board in the
873 investigation and conduct of a proceeding for licensure
874 revocation, suspension or restriction, including, but not limited
875 to, the cost of process service, court reporters, expert witnesses
876 and investigators.

877 Money collected by the board under paragraph (d)(iii) of this
878 section, shall be deposited to the credit of the Special Fund of
879 the Pharmacy Board;

880 (iv) The board may impose a monetary penalty for
881 those facilities/businesses registered with the Pharmacy Board as
882 wholesalers/manufacturers of not less than Three Hundred Dollars



883 (\$300.00) per violation and not more than Fifty Thousand Dollars
884 (\$50,000.00) per violation;

885 (v) The board may impose a monetary penalty for
886 any dispenser, pharmacist or practitioner licensed to dispense
887 controlled substance and specified noncontrolled substance drugs,
888 who knowingly fails to submit drug monitoring information or
889 knowingly submits incorrect dispensing information of not more
890 than Ten Thousand Dollars (\$10,000.00) per violation. Any penalty
891 collected under this paragraph (v) shall be deposited into the
892 special fund of the State Pharmacy Board to support the operations
893 of the Prescription Monitoring Program (PMP);

894 (vi) The board may impose a monetary penalty
895 for * * * any person * * * who obtains prescription information
896 and who knowingly discloses this information for misuse or
897 purposely alters the reporting information, or uses the PMP in any
898 manner other than for which it was intended, of not more * * *
899 than Fifty Thousand Dollars (\$50,000.00) per violation. Any
900 penalty collected under this paragraph (vi) shall be deposited
901 into the special fund of the State Board of Pharmacy and used to
902 support the operations of the Prescription Monitoring Program;

903 (vii) The board may impose a monetary penalty of
904 not more than One Thousand Dollars (\$1,000.00) per day upon any
905 person or business that practices or does business without the
906 license, registration or permit required by this chapter.



907 (e) Refusal to renew offender's license, registration
908 and/or permit;

909 (f) Placement of the offender on probation and
910 supervision by the board for a period to be determined by the
911 board;

912 (g) Public or private reprimand.

913 Whenever the board imposes any penalty under this subsection,
914 the board may require rehabilitation and/or additional education
915 as the board may deem proper under the circumstances, in addition
916 to the penalty imposed.

917 (2) Any person whose license, registration and/or permit has
918 been suspended, revoked or restricted pursuant to this chapter,
919 whether voluntarily or by action of the board, shall have the
920 right to petition the board at reasonable intervals for
921 reinstatement of such license, registration and/or permit. Such
922 petition shall be made in writing and in the form prescribed by
923 the board. Upon investigation and hearing, the board may, in its
924 discretion, grant or deny such petition, or it may modify its
925 original finding to reflect any circumstances which have changed
926 sufficiently to warrant such modifications. The procedure for the
927 reinstatement of a license, registration or permit that is
928 suspended for being out of compliance with an order for support,
929 as defined in Section 93-11-153, shall be governed by Section
930 93-11-157 or 93-11-163, as the case may be.



931 (3) Nothing herein shall be construed as barring criminal
932 prosecutions for violation of this chapter where such violations
933 are deemed as criminal offenses in other statutes of this state or
934 of the United States.

935 (4) A monetary penalty assessed and levied under this
936 section shall be paid to the board by the licensee, registrant or
937 permit holder upon the expiration of the period allowed for appeal
938 of such penalties under Section 73-21-101, or may be paid sooner
939 if the licensee, registrant or permit holder elects.

940 (5) When payment of a monetary penalty assessed and levied
941 by the board against a licensee, registrant or permit holder in
942 accordance with this section is not paid by the licensee,
943 registrant or permit holder when due under this section, the board
944 shall have the power to institute and maintain proceedings in its
945 name for enforcement of payment in the chancery court of the
946 county and judicial district of residence of the licensee,
947 registrant or permit holder, or if the licensee, registrant or
948 permit holder is a nonresident of the State of Mississippi, in the
949 Chancery Court of the First Judicial District of Hinds County,
950 Mississippi. When such proceedings are instituted, the board
951 shall certify the record of its proceedings, together with all
952 documents and evidence, to the chancery court and the matter shall
953 thereupon be heard in due course by the court, which shall review
954 the record and make its determination thereon. The hearing on the



955 matter may, in the discretion of the chancellor, be tried in
956 vacation.

957 (6) The board shall develop and implement a uniform penalty
958 policy which shall set the minimum and maximum penalty for any
959 given violation of board regulations and laws governing the
960 practice of pharmacy. The board shall adhere to its uniform
961 penalty policy except in such cases where the board specifically
962 finds, by majority vote, that a penalty in excess of, or less
963 than, the uniform penalty is appropriate. Such vote shall be
964 reflected in the minutes of the board and shall not be imposed
965 unless such appears as having been adopted by the board.

966 **SECTION 19.** Section 73-21-105, Mississippi Code of 1972, is
967 reenacted and amended as follows:

968 73-21-105. (1) Every facility/business that engages in the
969 wholesale distribution of prescription drugs, to include without
970 limitation, manufacturing in this state, distribution into this
971 state, or selling or offering to sell in this state, or
972 distribution from or within this state, and every reverse
973 distributor located in or outside of this state that conducts
974 business with pharmacies in this state, shall register biennially
975 or annually, to be determined by the board, with the Mississippi
976 State Board of Pharmacy by applying for a permit on a form
977 supplied by the board and accompanied by a fee as set by
978 subsection (4) of this section. The Pharmacy Board shall by



979 regulation determine the classification of permit(s) that shall be
980 required.

981 (2) Every business/facility/pharmacy located in this state
982 that engages in or proposes to engage in the dispensing and
983 delivery of prescription drugs to consumers shall register with
984 the Mississippi State Board of Pharmacy by applying for a permit
985 on a form supplied by the board and accompanied by a fee as set by
986 subsection (4) of this section. The Pharmacy Board shall by
987 regulation determine the classification of permit(s) that shall be
988 required.

989 (3) The board shall establish by rule or regulation the
990 criteria which each business shall meet to qualify for a permit in
991 each classification. The board shall issue a permit to any
992 applicant who meets the criteria as established. The board may
993 issue various types of permits with varying restrictions to
994 businesses where the board deems it necessary by reason of the
995 type of activities conducted by the business requesting a permit.

996 (4) The board shall specify by rule or regulation the
997 registration procedures to be followed, including, but not limited
998 to, specification of forms for use in applying for such permits
999 and times, places and fees for filing such applications. However,
1000 the biennial fee for an original or renewal permit shall not
1001 exceed Five Hundred Dollars (\$500.00), and the annual fee for an
1002 original or renewal permit shall not exceed Two Hundred Fifty
1003 Dollars (\$250.00).



1004 (5) Applications for permits shall include the following
1005 information about the proposed business:

1006 (a) Ownership;

1007 (b) Location;

1008 (c) Identity of the responsible person or pharmacist
1009 licensed to practice in the state, who shall be the pharmacist in
1010 charge of the pharmacy, where one is required by this chapter, and
1011 such further information as the board may deem necessary.

1012 (6) Permits issued by the board pursuant to this section
1013 shall not be transferable or assignable.

1014 (7) The board shall specify by rule or regulation minimum
1015 standards for the responsibility in the conduct of any
1016 business/facility and/or pharmacy that has been issued a permit.
1017 The board is specifically authorized to require that the portion
1018 of the facility located in this state to which a pharmacy permit
1019 applies be operated only under the direct supervision of no less
1020 than one (1) pharmacist licensed to practice in this state, and to
1021 provide such other special requirements as deemed necessary.
1022 Nothing in this subsection shall be construed to prevent any
1023 person from owning a pharmacy.

1024 (8) All businesses permitted by the board shall report to
1025 the board the occurrence of any of the following changes:

1026 (a) Permanent closing;

1027 (b) Change of ownership, management, location or
1028 pharmacist in charge;



1029 (c) Any and all other matters and occurrences as the
1030 board may require by rule or regulation.

1031 (9) Disasters, accidents and emergencies which may affect
1032 the strength, purity or labeling of drugs, medications, devices or
1033 other materials used in the diagnosis or the treatment of injury,
1034 illness and disease shall be immediately reported to the board.

1035 (10) No business that is required to obtain a permit shall
1036 be operated until a permit has been issued for such business by
1037 the board. Any person, firm or corporation violating any of the
1038 provisions of this section shall be guilty of a misdemeanor and,
1039 upon conviction thereof, shall be punished by a fine of not less
1040 than One Hundred Dollars (\$100.00) nor more than One Thousand
1041 Dollars (\$1,000.00), or imprisonment in the county jail for not
1042 less than thirty (30) days nor more than ninety (90) days, or by
1043 both such fine and imprisonment. However, the provisions of this
1044 chapter shall not apply to physicians, dentists, veterinarians,
1045 osteopaths or other practitioners of the healing arts who are
1046 licensed under the laws of the State of Mississippi and are
1047 authorized to dispense and administer prescription drugs in the
1048 course of their professional practice.

1049 **SECTION 20.** Section 73-21-106, Mississippi Code of 1972, is
1050 reenacted and amended as follows:

1051 73-21-106. (1) Any pharmacy located outside this state that
1052 ships, mails or delivers, in any manner, controlled substances or
1053 prescription or legend drugs or devices into this state shall be



1054 considered a nonresident pharmacy * * * and shall be permitted by
1055 the board * * *. The board shall establish by rule or regulation
1056 the criteria that each nonresident pharmacy must meet to qualify
1057 for a nonresident permit. After a permit has been issued, it may
1058 not be amended, transferred or reassigned. A pharmacist-in-charge
1059 of a nonresident pharmacy may not be the pharmacist-in-charge at
1060 any other location that has been issued a permit by the board.

1061 (2) * * * Each nonresident pharmacy shall:

1062 (a) * * * Comply with all lawful directions and
1063 requests for information from the regulatory or licensing agency
1064 of the state in which it is licensed as well as with all requests
1065 for information made by the board under this section. The
1066 nonresident pharmacy shall maintain at all times a valid unexpired
1067 license, permit or registration to conduct the pharmacy in
1068 compliance with the laws of the state in which it is a resident.
1069 As a prerequisite to being permitted by the board, the nonresident
1070 pharmacy shall submit a copy of the most recent inspection report
1071 resulting from an inspection conducted by the regulatory or
1072 licensing agency of the state in which it is located;

1073 (* * * b) Maintain its records of controlled substances
1074 and prescription or legend drugs or devices dispensed to patients
1075 in this state so that the records are readily retrievable from the
1076 records of other drugs dispensed; and

1077 (* * * c) Certify that it understands Mississippi
1078 pharmacy laws and regulations and agrees to comply with those laws



1079 and regulations and any other state or federal laws that apply to
1080 the practice of pharmacy. The pharmacist-in-charge must hold a
1081 Mississippi pharmacist license, be licensed to practice pharmacy
1082 in the state of residence of the nonresident pharmacy, and be
1083 current and in good standing with the licensing boards of both
1084 states.

1085 (* * *3) Any pharmacy subject to this section shall provide
1086 during its regular hours of operation, but not less than six (6)
1087 days per week and for a minimum of forty (40) hours per week, a
1088 toll-free telephone service to facilitate communication between
1089 patients in this state and a pharmacist at the pharmacy who has
1090 access to the patient's records. This toll-free number shall be
1091 disclosed on a label affixed to each container of drugs dispensed
1092 to patients in this state.

1093 (* * *4) The permit fee for nonresident pharmacies shall be
1094 the same as the fee as set by subsection (4) of Section 73-21-105.

1095 (* * *5) The permit requirements of this section shall
1096 apply to any nonresident pharmacy that dispenses, distributes,
1097 ships, mails or delivers controlled substances or prescription or
1098 legend drugs and devices into this state directly to a consumer.

1099 (* * *6) The board may deny, revoke or suspend a
1100 nonresident pharmacy permit only for:

1101 (a) Failure to comply with any requirement of this
1102 section or Section 41-29-125;



1103 (b) Conduct that causes serious bodily or serious
1104 psychological injury to a resident of this state if the board has
1105 referred the matter to the regulatory or licensing agency in the
1106 state in which the pharmacy is located and the regulatory or
1107 licensing agency fails to initiate an investigation within
1108 forty-five (45) days of the referral; or

1109 (c) Violation of the Uniform Controlled Substances Law.

1110 (* * *7) It is unlawful for any nonresident pharmacy that
1111 is not permitted under this section to advertise its services in
1112 this state, or for any person who is a resident of this state to
1113 advertise the pharmacy services of a nonresident pharmacy that is
1114 not permitted with the board, with the knowledge that the
1115 advertisement will or is likely to induce members of the public in
1116 this state to use the pharmacy to fill prescriptions.

1117 (* * *8) When requested to do so by the board or the
1118 Mississippi Bureau of Narcotics, each nonresident pharmacy shall
1119 supply any inspection reports, controlled substances dispensing
1120 records, warning notices, notice of deficiency reports or any
1121 other related reports from the state in which it is located
1122 concerning the operation of a nonresident pharmacy for review of
1123 compliance with state and federal drug laws.

1124 **SECTION 21.** Section 73-21-107, Mississippi Code of 1972, is
1125 reenacted as follows:

1126 73-21-107. (1) The board or its representative may enter
1127 and inspect, during reasonable hours, a facility which has



1128 obtained or applied for a permit under Section 73-21-105 relative
1129 to the following:

- 1130 (a) Drug storage and security;
- 1131 (b) Equipment;
- 1132 (c) Sanitary conditions; or
- 1133 (d) Records, reports, or other documents required to be
1134 kept or made under this chapter or the Uniform Controlled
1135 Substances Law (Section 41-29-101 et seq.) or rules and
1136 regulations adopted under such laws.

1137 (2) Prior to an entry and inspection, the board
1138 representative shall state his purpose and present appropriate
1139 credentials to the owner, pharmacist or agent in charge of a
1140 facility.

1141 (3) The board representative may:

1142 (a) Inspect and copy records, reports, and other
1143 documents required to be kept or made under this chapter, the
1144 Uniform Controlled Substances Law, or rules and regulations
1145 adopted under such laws;

1146 (b) Inspect, within reasonable limits and in a
1147 reasonable manner, a facility's storage, equipment, security,
1148 records, or prescription drugs or devices; or

1149 (c) Inventory any stock of any prescription drugs or
1150 devices in the facility.



1151 (4) Unless the owner, pharmacist, or agent in charge of the
1152 facility consents in writing, an inspection authorized by this
1153 section may not extend to:

1154 (a) Financial data;

1155 (b) Sales data other than shipment data; or

1156 (c) Pricing data.

1157 **SECTION 22.** Section 73-21-108, Mississippi Code of 1972, is
1158 reenacted as follows:

1159 73-21-108. (1) **Definitions.** For the purposes of this
1160 section:

1161 (a) "Home medical equipment" means technologically
1162 sophisticated medical equipment and devices usable in a home care
1163 setting, including, but not limited to:

1164 (i) Oxygen for human consumption, oxygen
1165 concentrators and/or oxygen delivery systems and equipment;

1166 (ii) Ventilators;

1167 (iii) Respiratory disease management devices;

1168 (iv) Electronic and computer driven wheelchairs
1169 and seating systems;

1170 (v) Apnea monitors;

1171 (vi) Transcutaneous electrical nerve stimulator
1172 (TENS) units;

1173 (vii) Low air loss cutaneous pressure management
1174 devices;

1175 (viii) Sequential compression devices;



1176 (ix) Neonatal home phototherapy devices;
1177 (x) Feeding pumps; and
1178 (xi) Other similar equipment as defined in
1179 regulations adopted by the board.

1180 The term "home medical equipment" does not include medical
1181 equipment used in the normal course of treating patients by
1182 hospitals, hospices, long-term care facilities or home health
1183 agencies, or medical equipment used or dispensed by health care
1184 professionals licensed by the State of Mississippi if the
1185 professional is practicing within the scope of his or her
1186 professional practice. In addition, the term does not include
1187 items such as upper and lower extremity prosthetics, canes,
1188 crutches, walkers, bathtub grab bars, standard wheelchairs,
1189 commode chairs and bath benches.

1190 (b) "Home medical equipment services" means the
1191 delivery, installation, maintenance, replacement, and/or
1192 instruction in the use of home medical equipment, used by a sick
1193 or disabled individual, to allow the individual to be cared for
1194 and maintained in a home or noninstitutional environment.

1195 (c) "Medical gas" means those gases and liquid oxygen
1196 intended for human consumption.

1197 (d) "Order" means an order issued by a licensed
1198 practitioner legally authorized to order home medical equipment
1199 and/or medical gases.



1200 (2) **Permit required.** (a) No person, business or entity
1201 located in this state or outside of this state that is subject to
1202 this section shall sell, rent or provide or offer to sell, rent or
1203 provide directly to patients in this state any home medical
1204 equipment, legend devices, and/or medical gas unless such person,
1205 business or entity first obtains a Medical Equipment Supplier
1206 Permit from the board.

1207 (b) The permitting requirements of this section apply
1208 to all persons, companies, agencies and other business entities
1209 that are in the business of supplying home medical equipment to
1210 patients in their places of residence and that bill the patient or
1211 the patient's insurance, Medicare, Medicaid or other third party
1212 payor for the rent or sale of that equipment.

1213 (c) The board shall require a separate permit for each
1214 facility location directly or indirectly owned or operated in this
1215 state.

1216 (d) The application for a permit shall be made to the
1217 board on a form supplied by the board and shall be accompanied by
1218 a fee of not more than Three Hundred Dollars (\$300.00), as
1219 prescribed by the board. Once issued, every permit must be
1220 renewed annually, and the renewal fee shall be not more than One
1221 Hundred Seventy-five Dollars (\$175.00), as prescribed by the
1222 board.

1223 (e) All permits issued under this section shall expire
1224 annually on June 30 of each year. Applications for renewal must



1225 be made to the board on or before June 30 and must be accompanied
1226 by the fee as prescribed by the board. A late renewal fee of One
1227 Hundred Dollars (\$100.00) shall be added to all renewal
1228 applications received by the board after June 30 of each renewal
1229 period. The permit shall become void if the renewal application,
1230 renewal fee and the late renewal fee are not received by the board
1231 by September 30 of each year.

1232 (3) **Exemptions.** (a) The permitting requirements of this
1233 section do not apply to the following entities or practitioners
1234 unless they have a separate business entity, company, corporation
1235 or division that is in the business of providing home medical
1236 equipment for sale or rent to patients at their places of
1237 residence:

- 1238 (i) Home health agencies;
- 1239 (ii) Hospitals;
- 1240 (iii) Wholesalers and/or manufacturers;
- 1241 (iv) Medical doctors, physical therapists,
1242 respiratory therapists, occupational therapists, speech
1243 pathologists, optometrists, chiropractors and podiatrists who use
1244 home medical equipment and/or legend devices in their individual
1245 practices;
- 1246 (v) Pharmacies;
- 1247 (vi) Hospice programs;
- 1248 (vii) Nursing homes and/or long-term care
1249 facilities;



1250 (viii) Veterinarians; dentists; and emergency
1251 medical services.

1252 (b) Although community pharmacies are exempt from the
1253 permitting requirements of this section, they shall be subject to
1254 the same regulations that are applicable to permitted businesses
1255 or entities for the sale or rental of home medical equipment
1256 covered by this section.

1257 (c) Nothing in this section shall prohibit trained
1258 individuals from using oxygen, liquid oxygen and/or legend devices
1259 in emergencies.

1260 (d) Nothing in this section shall prohibit the
1261 prehospital emergency administration of oxygen by licensed health
1262 care providers, emergency medical technicians, first responders,
1263 fire fighters, law enforcement officers and other emergency
1264 personnel trained in the proper use of emergency oxygen.

1265 (4) **Order required.** Home medical equipment suppliers shall
1266 not provide any home medical equipment to a patient without a
1267 valid order from an authorized licensed practitioner.

1268 (5) **Regulations.** The board shall adopt regulations for the
1269 distribution and sale or rental of home medical equipment, legend
1270 devices and medical gases that promote the public health and
1271 welfare and comply with at least the minimum standards, terms and
1272 conditions of federal laws and regulations. The regulations shall
1273 include, without limitation:



1274 (a) Minimum information from each home medical
1275 equipment, legend device and medical gas supplier required for
1276 permitting and renewal permits;

1277 (b) Minimum qualifications of persons who engage in the
1278 distribution of home medical equipment;

1279 (c) Appropriate education, training or experience of
1280 persons employed by home medical equipment suppliers;

1281 (d) Minimum standards for storage of home medical
1282 equipment;

1283 (e) Minimum requirements for the establishment and
1284 maintenance of all records for the sale, rental and servicing of
1285 home medical equipment; and

1286 (f) Minimum standards of operation and professional
1287 conduct.

1288 (6) **Medical Equipment Advisory Committee to the board.**

1289 (a) A Medical Equipment Advisory Committee (MEAC),
1290 composed of three (3) members selected by the Mississippi
1291 Association of Medical Equipment Suppliers and approved by the
1292 board, shall review and make recommendations to the board
1293 regarding all regulations dealing with home medical equipment,
1294 legend devices and medical gases that are proposed by the board
1295 and before they are adopted by the board.

1296 (b) All MEAC members must have been actively involved
1297 in the home medical equipment business for a minimum of five (5)
1298 years before the selection to the committee and shall hold and



1299 maintain, in good standing, a permit issued by the board under
1300 this section.

1301 (c) The MEAC members shall meet at least quarterly and
1302 review all home medical equipment suppliers' inspection reports.
1303 All complaints and reports of investigations of violations of law
1304 or regulations regarding home medical equipment, legend devices
1305 and medical gases shall first be reviewed by the MEAC. After
1306 review, the MEAC may make recommendations to the board's
1307 Investigations Review Committee regarding further administrative
1308 action by the board.

1309 (d) The MEAC shall keep and maintain minutes of all
1310 meetings of the MEAC and shall provide copies of the minutes to
1311 the board on a quarterly basis.

1312 (7) **Revocation, suspension or restriction of permit and**
1313 **penalties.**

1314 (a) The board may revoke, suspend, restrict or refuse
1315 to issue or renew a permit or impose a monetary penalty, in
1316 accordance with Section 73-21-103 except that the monetary penalty
1317 shall not exceed Ten Thousand Dollars (\$10,000.00) per violation,
1318 if the business or holder of a permit or applicant for a permit
1319 issued under this section has committed or is found guilty by the
1320 board of any of the following:

1321 (i) Violation of any federal, state or local law
1322 or regulations relating to home medical equipment, legend devices
1323 or medical gases.



1324 (ii) Violation of any of the provisions of this
1325 section or regulations adopted under this section.

1326 (iii) Commission of an act or engaging in a course
1327 of conduct that constitutes a clear and present danger to the
1328 public health and safety.

1329 (iv) Filing a claim or assisting in the filing of
1330 a claim for reimbursement for home medical equipment or home
1331 medical equipment services that were not provided or that were not
1332 authorized to be provided.

1333 (v) Failure to comply with any lawful order of the
1334 board.

1335 (b) Disciplinary action by the board against a business
1336 or any person holding a permit under this section shall be in
1337 accordance with Section 73-21-99.

1338 **SECTION 23.** Section 73-21-109, Mississippi Code of 1972, is
1339 reenacted as follows:

1340 73-21-109. No person shall make use of the terms
1341 "drugstore," "pharmacy," "apothecary" or words of similar meaning
1342 which indicate that pharmaceutical services are performed in any
1343 sign, letterhead or advertisement unless such person is a permit
1344 holder as provided in Section 73-21-105, or such property or name
1345 was previously registered with the Mississippi State Board of
1346 Pharmacy or provided pharmaceutical services in excess of twenty
1347 (20) years. Any person violating this section shall be guilty of
1348 a misdemeanor and, upon conviction thereof, shall be punished by a



1349 fine of not less than One Hundred Dollars (\$100.00) nor more than
1350 Three Hundred Dollars (\$300.00), or by imprisonment in the county
1351 jail for not less than thirty (30) days nor more than ninety (90)
1352 days, or by both.

1353 **SECTION 24.** Section 73-21-111, Mississippi Code of 1972, is
1354 reenacted as follows:

1355 73-21-111. (1) The board shall make, adopt, amend and
1356 repeal from time to time such rules and regulations for the
1357 regulation of supportive personnel as may be deemed necessary by
1358 the board.

1359 (2) Every person who acts or serves as a pharmacy technician
1360 in a pharmacy that is located in this state and permitted by the
1361 board shall obtain a registration from the board. To obtain a
1362 pharmacy technician registration the applicant must:

1363 (a) Have submitted a written application on a form(s)
1364 prescribed by the board; and

1365 (b) Be of good moral character; and

1366 (c) Have paid the initial registration fee not to
1367 exceed One Hundred Dollars (\$100.00).

1368 (3) Each pharmacy technician shall renew his or her
1369 registration annually. To renew his or her registration, a
1370 technician must:

1371 (a) Submit an application on a form prescribed by the
1372 board; and



1373 (b) Pay a renewal fee not to exceed One Hundred Dollars
1374 (\$100.00) for each annual registration period. The board may add
1375 a surcharge of not more than Five Dollars (\$5.00) to the
1376 registration renewal fee to assist in funding a program that
1377 assists impaired pharmacists, pharmacy students and pharmacy
1378 technicians.

1379 (4) To insure that all applicants are of good moral
1380 character, the board shall conduct a criminal history records
1381 check on all applicants for a license. In order to determine the
1382 applicant's suitability for licensing, the applicant shall be
1383 fingerprinted. The board shall submit the fingerprints to the
1384 Department of Public Safety for a check of the state criminal
1385 records and forwarded to the Federal Bureau of Investigation for a
1386 check of the national criminal records. The Department of Public
1387 Safety shall disseminate the results of the state check and the
1388 national check to the board for a suitability determination. The
1389 board shall be authorized to collect from the applicant the amount
1390 of the fee that the Department of Public Safety charges the board
1391 for the fingerprinting, whether manual or electronic, and the
1392 state and national criminal history records checks.

1393 **SECTION 25.** Section 73-21-113, Mississippi Code of 1972, is
1394 reenacted as follows:

1395 73-21-113. All fees received by the board from examinations,
1396 licenses, permits and monetary penalties, and any other funds
1397 received by the board, shall be paid to the State Treasurer, who



1398 shall issue receipts therefor and deposit such funds in the State
1399 Treasury in a special fund to the credit of the board. All such
1400 funds shall be expended only pursuant to appropriation approved by
1401 the Legislature and as provided by law.

1402 **SECTION 26.** Section 73-21-115, Mississippi Code of 1972, is
1403 reenacted as follows:

1404 73-21-115. (1) Every prescription written in this state by
1405 a person authorized to issue such prescription shall be on
1406 prescription forms containing two (2) lines for the prescriber's
1407 signature. There shall be a signature line in the lower
1408 right-hand corner of the prescription form beneath which shall be
1409 clearly imprinted the words "substitution permissible." There
1410 shall be a signature line in the lower left-hand corner of the
1411 prescription form beneath which shall be clearly imprinted the
1412 words "dispense as written." The prescriber's signature on either
1413 signature line shall validate the prescription and shall designate
1414 approval or disapproval of product selection.

1415 (2) If a prescription form which does not contain the two
1416 (2) signature lines required in subsection (1) of this section is
1417 utilized by the prescriber, he shall write in his own handwriting
1418 the words "dispense as written" thereupon to prevent product
1419 selection.

1420 (3) A pharmacist licensed by the Mississippi State Board of
1421 Pharmacy may dispense a one-time emergency dispensing of a
1422 prescription of up to a seventy-two-hour supply of a prescribed



1423 medication in the event the pharmacist is unable to contact the
1424 prescriber to obtain refill authorization, provided that:

1425 (a) The prescription is not for a controlled substance;

1426 (b) In the pharmacist's professional judgment, the
1427 interruption of therapy might reasonably produce undesirable
1428 health consequences or may cause physical or mental discomfort;

1429 (c) The dispensing pharmacist notifies the prescriber
1430 or his agent of the emergency dispensing within seven (7) working
1431 days after the one-time emergency dispensing;

1432 (d) The pharmacist properly records the dispensing as a
1433 separate nonrefillable prescription. Said document shall be filed
1434 as is required of all other prescription records. This document
1435 shall be serially numbered and contain all information required of
1436 other prescriptions. In addition it shall contain the number of
1437 the prescription from which it was refilled; and

1438 (e) The pharmacist shall record on the new document the
1439 circumstances which warrant this emergency dispensing.

1440 This emergency dispensing shall be done only in the permitted
1441 facility which contains the nonrefillable prescription.

1442 **SECTION 27.** Section 73-21-117, Mississippi Code of 1972, is
1443 reenacted as follows:

1444 73-21-117. (1) A pharmacist may select a generic equivalent
1445 drug product only when such selection results in lower cost to the
1446 purchaser, unless product selection is expressly prohibited by the
1447 prescriber.



1448 (2) A pharmacist shall select a generic equivalent drug
1449 product when:

1450 (a) The purchaser requests the selection of a generic
1451 equivalent drug product;

1452 (b) The prescriber has not expressly prohibited product
1453 selection; and

1454 (c) Product selection will result in lower cost to the
1455 purchaser.

1456 Before product selection is made, the pharmacist shall advise
1457 the purchaser of his prerogatives under this subsection.

1458 (3) When requested by the purchaser to dispense the drug
1459 product as ordered by the prescriber, a pharmacist shall not
1460 select a generic equivalent drug product.

1461 **SECTION 28.** Section 73-21-119, Mississippi Code of 1972, is
1462 reenacted as follows:

1463 73-21-119. (1) The label of the container of any drug
1464 product which is sold within the State of Mississippi for resale
1465 at retail and which requires a prescription to be dispensed at
1466 retail shall contain at a minimum the name of the manufacturer of
1467 the final dosage unit, expiration date if applicable, batch or lot
1468 number and national drug code.

1469 (2) Whenever product selection is made, the pharmacist shall
1470 indicate on the label of the dispensed container the initials
1471 "G.E." and the proprietary name of the product dispensed or the
1472 generic name of the product dispensed and its manufacturer either



1473 written in full or appropriately abbreviated, unless the
1474 prescriber indicates that the name of the drug product shall not
1475 appear on the label.

1476 **SECTION 29.** Section 73-21-121, Mississippi Code of 1972, is
1477 reenacted as follows:

1478 73-21-121. (1) Product selection as authorized by Sections
1479 73-21-115 through 73-21-119 shall not constitute evidence of
1480 negligence by the dispensing pharmacist when such product
1481 selection is in accordance with reasonable and prudent pharmacy
1482 practice. No prescriber shall be liable for civil damages or in
1483 any criminal prosecution arising from the incorrect product
1484 selection by a pharmacist.

1485 (2) Any person having knowledge relating to a pharmacist or
1486 to a pharmacy student which might provide grounds for disciplinary
1487 action by the board may report relevant facts to the board, and
1488 shall by reason of reporting such facts in good faith be immune
1489 from civil liability.

1490 (3) Any person furnishing information in the form of data,
1491 reports or records to the board or to a pharmacist organization
1492 approved by the board to receive such information, where such
1493 information is furnished for the purpose of aiding a pharmacist or
1494 a pharmacy student impaired by chemical abuse or by mental or by
1495 physical illness, shall by reason of furnishing such information
1496 in good faith be immune from civil liability.



1497 (4) The records of the board or the records of a pharmacist
1498 organization approved by the board to aid pharmacists or pharmacy
1499 students impaired by chemical abuse, where such records relate to
1500 the impairment, shall be confidential and are not considered open
1501 records; provided, however, the board may disclose this
1502 confidential information only:

1503 (a) In a disciplinary hearing before the board, or in
1504 an appeal of an action or order of the board;

1505 (b) To the pharmacist licensing or disciplinary
1506 authorities of other jurisdictions in the case of a pharmacist who
1507 is licensed in, or seeking transfer to, another state; or

1508 (c) Pursuant to an order of a court of competent
1509 jurisdiction.

1510 **SECTION 30.** Section 73-21-123, Mississippi Code of 1972, is
1511 reenacted as follows:

1512 73-21-123. Nothing in this chapter shall be construed to
1513 prevent, or in any manner interfere with, or to require a permit
1514 for the sale of nonnarcotic nonprescription drugs which may be
1515 lawfully sold under the United States Food, Drug and Cosmetic Act
1516 (21 USCS 301 et seq. as now or hereafter amended) without a
1517 prescription, nor shall any rule or regulation be adopted by the
1518 board under the provisions of this chapter which shall require the
1519 sale of nonprescription drugs by a licensed pharmacist of in a
1520 pharmacy or otherwise apply to or interfere with the sale or
1521 distribution of such drugs.



1522 **SECTION 31.** Section 73-21-125, Mississippi Code of 1972, is
1523 brought forward as follows:

1524 73-21-125. (1) Any community pharmacy, including a
1525 faith-based community pharmacy, or any licensed pharmacist who
1526 voluntarily provides charitable services in a community pharmacy,
1527 or any other person who serves as a volunteer in a community
1528 pharmacy, shall be immune from liability for any civil action
1529 arising out of supplying pharmaceutical products in the course of
1530 providing such charitable or gratuitous pharmaceutical products.
1531 This section shall not extend immunity to acts of gross negligence
1532 or willful or wanton misconduct or to the manufacturer or designer
1533 of products provided.

1534 (2) Any community pharmacy seeking immunity under this
1535 section shall post a notice, in a conspicuous place adjacent to
1536 the area where prescriptions are picked up by consumers, reading
1537 substantially as follows: "NOTICE: If you are harmed by
1538 medication that you receive here, you do not have the same legal
1539 recourse as you have against other pharmacies." Failure to post
1540 the notice negates the immunity from liability provided under this
1541 section. The notice shall be no less than eleven (11) by fourteen
1542 (14) inches in size, and the type used shall be no smaller than
1543 thirty-six (36) point and surrounded by a one-inch solid black
1544 border.

1545 (3) For purposes of this section, "community pharmacy" means
1546 a pharmacy operated solely for charitable purposes, whose only



1547 function is to supply gratuitous pharmaceutical products, and
1548 which is operated by a nonprofit organization qualified or
1549 eligible for qualification as a tax-exempt organization under 26
1550 USCS 501.

1551 **SECTION 32.** Section 73-21-126, Mississippi Code of 1972, is
1552 amended as follows:

1553 73-21-126. (1) The State Board of Pharmacy shall promulgate
1554 rules regarding the issuance and renewal of licenses and permits
1555 for new or renewal application requirements for both in_ and
1556 out_of_state wholesale distributors, chain pharmacy warehouses and
1557 repackagers shipping into Mississippi. Requirements for new
1558 and * * */or renewal applications, if information has not been
1559 previously provided to the board, will include, but not be limited
1560 to, the following:

1561 (a) Type of ownership (individual, partnership or
1562 corporation);

1563 (b) Names of principal owners or officers and social
1564 security numbers;

1565 (c) Names of designated representatives and social
1566 security numbers;

1567 (d) Criminal background checks of applicants and
1568 designated representatives as required by rule;

1569 (e) Copy of license in home state;

1570 (f) Bond requirements.



1571 (2) The board shall promulgate rules for the establishment
1572 of a pedigree or electronic file to be used by wholesale
1573 distributors, chain pharmacy warehouses and repackagers for the
1574 purpose of ensuring the integrity of drugs owned, purchased,
1575 distributed, returned, transferred and sold when the products
1576 leave the normal distribution channel.

1577 (3) The board is authorized to use an outside agency to
1578 accredit wholesale distributors and repackagers, including the
1579 National Association of Boards of Pharmacy's (NABP) Verified
1580 Accredited Wholesale Distributors (VAWD) program.

1581 (4) Pharmacies shall not be responsible for verification or
1582 adjudication of the pedigree for pharmaceuticals.

1583 (5) The board may exempt wholesalers accredited by the VAWD
1584 program from the above requirements.

1585 **SECTION 33.** Section 73-21-127, Mississippi Code of 1972, is
1586 amended as follows:

1587 73-21-127. The Board of Pharmacy shall develop and implement
1588 a computerized program to track prescriptions for controlled
1589 substances and to report suspected abuse and misuse of controlled
1590 substances in compliance with the federal regulations promulgated
1591 under authority of the National All Schedules Prescription
1592 Electronic Reporting Act of 2005 and in compliance with the
1593 federal HIPAA law, under the following conditions:

1594 (a) Submission or reporting of dispensing information
1595 shall be mandatory and required by the State Board of Pharmacy for



1596 any entity dispensing controlled substances in or into the State
1597 of Mississippi, except for the dispensing of controlled substance
1598 drugs * * * by a veterinarian residing in the State of
1599 Mississippi.

1600 (b) The prescriptions tracked shall be prescriptions
1601 for controlled substances listed in * * * Schedule II, III, IV or
1602 V and specified noncontrolled substances * * * identified by the
1603 State Board of Pharmacy that are dispensed to residents in the
1604 State of Mississippi by licensed pharmacies, nonresident
1605 pharmacies, institutions and dispensing practitioners, regardless
1606 of dispenser location.

1607 (c) The Board of Pharmacy shall report any activity it
1608 reasonably suspects may be fraudulent or illegal to the
1609 appropriate law enforcement agency or occupational licensing board
1610 and provide them with the relevant information obtained for
1611 further investigation.

1612 (d) The program shall provide information regarding the
1613 potential inappropriate use of controlled substances and the
1614 specified noncontrolled substances to practitioners,
1615 pharmacists-in-charge and appropriate state agencies in order to
1616 prevent the inappropriate or illegal use of these controlled
1617 substances. The specific purposes of the program shall be to: be
1618 proactive in safeguarding public health and safety; support the
1619 legitimate use of controlled substances; facilitate and encourage
1620 the identification, intervention with and treatment of individuals



1621 addicted to controlled substances and specified noncontrolled
1622 drugs; identify and prevent drug diversion; provide assistance to
1623 those state and federal law enforcement and regulatory agencies
1624 investigating cases of drug diversion or other misuse; and inform
1625 the public and health care professionals of the use and abuse
1626 trends related to controlled substance and specified noncontrolled
1627 drugs.

1628 (e) (i) Access to collected data shall be confidential
1629 and not subject to the provisions of the federal Freedom of
1630 Information Act or the Mississippi Open Records Act. Upon
1631 request, the State Board of Pharmacy shall provide collected
1632 information to: pharmacists or practitioners who are properly
1633 registered with the State Board of Pharmacy and are authorized to
1634 prescribe or dispense controlled substances for the purpose of
1635 providing medical and pharmaceutical care for their patients;
1636 local, state and federal law enforcement officials engaged in the
1637 administration, investigation or enforcement of the laws governing
1638 illicit drug use; regulatory and licensing boards in this state;
1639 Division of Medicaid regarding Medicaid and Medicare Program
1640 recipients; judicial authorities under grand jury subpoena; an
1641 individual who requests the individual's own prescription
1642 monitoring information; and prescription monitoring programs in
1643 other states through mutual agreement adhering to State Board of
1644 Pharmacy policies.



1645 (ii) The Director of the Mississippi Bureau of
1646 Narcotics, or his designee, shall have access to the Prescription
1647 Monitoring Program (PMP) database for the purpose of investigating
1648 the potential illegal acquisition, distribution, dispensing,
1649 prescribing or administering of the controlled and noncontrolled
1650 substances monitored by the program, subject to all legal
1651 restrictions on further dissemination of the information obtained.

1652 (iii) The State Board of Pharmacy may also provide
1653 * * * statistical data for research or educational purposes if
1654 the board determines the use of the data to be of significant
1655 benefit to public health and safety. The board maintains the
1656 right to refuse any request for PMP data.

1657 (iv) A pharmacist licensed by the Mississippi
1658 Board of Pharmacy must be a registered user of the PMP. Failure
1659 of a pharmacist licensed by the Mississippi Board of Pharmacy to
1660 register as a user of the PMP is grounds for disciplinary action
1661 by the board.

1662 (f) The Prescription Monitoring Program through the
1663 Board of Pharmacy may:

1664 (i) Establish the cost of administration,
1665 maintenance, and operation of the program and charge to like
1666 agencies a fee based on a formula to be determined by the board
1667 with collaboration and input from participating agencies; and

1668 (ii) Assess charges for information and/or
1669 statistical data provided to agencies, institutions and



1670 individuals. The amounts of those fees shall be set by the
1671 Executive Director of the Board of Pharmacy based on the
1672 recommendation of the Director of the PMP.

1673 All such fees collected shall be deposited into the special
1674 fund of the State Board of Pharmacy and used to support the
1675 operations of the PMP.

1676 (* * *g) A dispenser pharmacist or practitioner
1677 licensed to dispense controlled substances and specified
1678 noncontrolled substance drugs who knowingly fails to submit drug
1679 monitoring information or knowingly submits incorrect dispensing
1680 information shall be subject to actions against the pharmacist's
1681 or practitioner's license, registrations or permit and/or an
1682 administrative penalty as provided in Sections 73-21-97 and
1683 73-21-103. Any misuse of the PMP is subject to penalties as
1684 provided in Sections 73-21-97 and 73-21-103.

1685 (h) The Board of Pharmacy and the Prescription
1686 Monitoring Program shall be immune from civil liability arising
1687 from inaccuracy of any of the information submitted to the
1688 program.

1689 (* * *i) "Practitioner," as used in this section,
1690 shall include any person licensed, registered or otherwise
1691 permitted to distribute, dispense, prescribe or administer a
1692 controlled substance, as defined under Section 41-29-105(y).

1693 (* * *j) In addition to any funds appropriated by the
1694 Legislature, the State Board of Pharmacy may apply for any



1695 available grants and accept any gifts, grants or donations to
1696 assist in future development or in maintaining the program.

1697 * * *

1698 **SECTION 34.** Section 73-21-129, Mississippi Code of 1972, is
1699 amended as follows:

1700 73-21-129. (1) Each manufacturer whose products are
1701 distributed within the State of Mississippi shall make adequate
1702 provision for the return of outdated drugs from pharmacies, both
1703 full and partial containers, excluding biological, infused or
1704 intravenously injected drugs and drugs that are inhaled during
1705 surgery, within six (6) months after the labeled expiration date,
1706 for prompt full credit or refund.

1707 (2) Wholesale distributors and reverse distributors that are
1708 required to register with the board and have a permit under
1709 Section 73-21-105 shall implement and administer the return
1710 policies established by the manufacturer.

1711 (3) If the board receives information that a manufacturer
1712 has failed to comply with this section, the board shall
1713 investigate the matter and present any evidence of the
1714 manufacturer's failure to comply to a review committee composed of
1715 the Dean of the University of Mississippi School of Pharmacy, the
1716 Executive Director of the State Board of Pharmacy and the Director
1717 of the Pharmacy Bureau of the Division of Medicaid, or the
1718 designee of any of those officials. The committee shall review
1719 the evidence of the manufacturer's failure to comply with this



1720 section and make a recommendation to the board regarding the
1721 discipline of the manufacturer for its failure to comply. After
1722 the board has received the recommendation of the committee, the
1723 board may discipline the manufacturer by providing that the
1724 manufacturer's products shall be ineligible for use in product
1725 selection in any state drug assistance programs.

1726 (4) A pharmacist may not dispense a prescription drug or
1727 controlled drug unless the pharmacist has satisfactory evidence
1728 that the manufacturer of the drug has a procedure for the return
1729 of expired drugs.

1730 (5) Any manufacturer that had a repurchase program in place
1731 on January 1, 2008, shall be exempt from the provisions of this
1732 section, provided that the repurchase program makes provision for
1733 the repurchase of outdated drugs in either full or partial amounts
1734 within six (6) months after the labeled expiration date.

1735 (6) As used in this section, the term "biological drug" or
1736 "biological product" means a virus, therapeutic serum, toxin,
1737 antitoxin, vaccine, blood, blood component or derivative,
1738 allergenic product or analogous product, or arsphenamine or
1739 derivative of arsphenamine or any other trivalent organic arsenic
1740 compound, applicable to the prevention, treatment or cure of a
1741 disease or condition of human beings.

1742 * * *

1743 **SECTION 35.** This act shall take effect and be in force from
1744 and after July 1, 2016.

