WEST VIRGINIA LEGISLATURE

REGULAR SESSION, 1995

ENROLLED

Com. Sub. for
HOUSE BILL No. 2451

(By Delegates Gallagher & Boden...)

Passed March 11, 1995

In Effect 90 Days from Passage
AN ACT to repeal sections twelve-a and sixteen-a, article five, chapter thirty of the code of West Virginia, one thousand nine hundred thirty-one, as amended; to amend and reenact sections one, two, three, four, five, six, seven, eight, nine, ten, eleven, twelve, twelve-b, thirteen, fourteen, fourteen-a, fifteen, sixteen, nineteen, twenty-one and twenty-two of said article; and to further amend said article five by adding thereto nine new sections, designated sections one-a, one-b, two-a, five-a, seven-a, seven-b, sixteen-b, sixteen-c and twenty-two-a, all relating to the regulation of pharmacists, licensed interns and pharmacist technicians; repealing existing section twelve-a relating to drug and drug price listing and posting requirements and penalties for noncompliance; repealing existing section sixteen-a authorizing the manufacture of laetrile; legislative findings and statement of purpose; defining terms; filling of board vacancies; board qualifications; increasing board compensation; meetings and business of the board; clarifying public and closed meetings; records kept by the board; providing for expungement of records; examination of records; notice requirements; public information; making various technical changes; permitting licensed interns and pharmacy technicians to assist pharmacists; experience and training qualifications for pharmacists, licensed interns and
pharmacy technicians; titles and terms; regulating pharmacy technicians; reciprocity; disciplinary proceedings; grounds for disciplinary action; fines and penalties; hearings and notice; confidentiality of prescription records; reporting criteria for professional malpractice, incompetence and convictions; voluntary agreements relating to alcohol or chemical dependency; confidentiality requirements; pharmacy lists; fees; license renewals and display; prohibitions; distribution of generic and brand-name drugs; prescription requirements for 'brand medically necessary' drugs; requiring ownership of USP-DI reference manual; pharmacy registration; pharmacists-in-charge; increasing fines for violations of equipment requirements; manufacturing permits; authorizing partial filling of schedule II medications under certain circumstances; limitations on application of article; increasing criminal and civil penalties; providing for immunity from civil actions for board members; limiting liability for professionals reporting to the board; required reporting of litigation results to the board; and rule-making authority.

Be it enacted by the Legislature of West Virginia:

That sections twelve-a and sixteen-a, article five, chapter thirty of the code of West Virginia, one thousand nine hundred thirty-one, as amended, be repealed; that sections one, two, three, four, five, six, seven, eight, nine, ten, eleven, twelve, twelve-b, thirteen, fourteen, fourteen-a, fifteen, sixteen, nineteen, twenty-one and twenty-two of said article be amended and reenacted; and that said article five be further amended by adding theretoeight new sections, designated sections one-a, one-b, two-a, five-a, seven-a, seven-b, sixteen-b, sixteen-c and twenty-two-a, all to read as follows:

ARTICLE 5. PHARMACISTS, PHARMACY TECHNICIANS, PHARMACY INTERNS AND PHARMACIES.

§30-5-1. Legislative findings.

The Legislature hereby finds and declares that the practice of pharmacy is a privilege and not a natural or fundamental right of any individual. As a matter of public policy, it is necessary to protect the public through the enactment of this article and to regulate the granting of such privileges and their use. This article shall be liberally
§30-5-la. Statement of purpose.

It is the purpose of this article to promote, preserve and protect the public health, safety and welfare by the effective regulation of the practice of pharmacy; the licensure of pharmacists; the licensure, and regulation of all sites or persons who distribute, manufacture, or sell drugs or devices used in the dispensing and administration of drugs or devices within this state.

§30-5-lb. Definitions.

The following words and phrases as used in this article, shall have the following meanings, unless the context otherwise requires:

(a) "Administer" means the direct application of a drug to the body of a patient or research subject by injection, inhalation, ingestion or any other means.

(b) "Board of pharmacy" or "board" means the West Virginia state board of pharmacy.

(c) "Compounding" means:

(1) The preparation, mixing, assembling, packaging or labeling of a drug or device:

(A) As the result of a practitioner's prescription drug order or initiative based on the practitioner/patient/pharmacist relationship in the course of professional practice for sale or dispensing; or

(B) For the purpose of, or as an incident to, research, teaching or chemical analysis and not for sale or dispensing;

(2) The preparation of drugs or devices in anticipation of prescription drug orders based on routine, regularly observed prescribing patterns.

(d) "Confidential information" means information maintained by the pharmacist in the patient record or which is communicated to the patient as part of patient counseling, or which is communicated by the patient to
the pharmacist. This information is privileged and may be
released only to the patient or to other members of the
health care team and other pharmacists where, in the
pharmacist's professional judgment, such release is neces-
sary to the patient's health and well-being; to such other
persons or governmental agencies authorized by law to
receive such privileged information; as necessary for the
limited purpose of peer review and utilization review; as
authorized by the patient or required by court order.

(e) "Deliver" or "delivery" means the actual, construc-
tive or attempted transfer of a drug or device from one
person to another, whether or not for a consideration.

(f) "Device" means an instrument, apparatus, imple-
ment or machine, contrivance, implant or other similar or
related article, including any component part or accessory,
which is required under federal law to bear the label, "Caution: Federal or state law requires dispensing by or on the
order of a physician."

(g) "Dispense" or "dispensing" means the preparation
and delivery of a drug or device in an appropriately la-
beled and suitable container to a patient or patient's repre-
sentative or surrogate pursuant to a lawful order of a prac-
titioner for subsequent administration to, or use by, a pa-
tient.

(h) "Distribute" means the delivery of a drug or device
other than by administering or dispensing.

(i) "Drug" means:

(1) Articles recognized as drugs in the USP-DI, Facts
and Comparisons, Physicians Desk Reference or supple-
ments thereto, for use in the diagnosis, cure, mitigation,
treatment or prevention of disease in human or other ani-
imals;

(2) Articles, other than food, intended to affect the
structure or any function of the body of human or other
animals; and

(3) Articles intended for use as a component of any
articles specified in subsection (1) or (2) of this section.
(j) "Drug regimen review" includes, but is not limited to, the following activities:

(1) Evaluation of the prescription drug orders and patient records for:
   (A) Known allergies;
   (B) Rational therapy-contraindications;
   (C) Reasonable dose and route of administration; and
   (D) Reasonable directions for use.

(2) Evaluation of the prescription drug orders and patient records for duplication of therapy.

(3) Evaluation of the prescription drug for interactions and/or adverse effects which may include, but are not limited to, any of the following:
   (A) Drug-drug;
   (B) Drug-food;
   (C) Drug-disease; and
   (D) Adverse drug reactions.

(4) Evaluation of the prescription drug orders and patient records for proper utilization, including over utilization and under utilization and optimum therapeutic outcomes.

(k) "Intern" means an individual who is:

(1) Currently registered by this state to engage in the practice of pharmacy while under the supervision of a licensed pharmacist and is satisfactorily progressing toward meeting the requirements for licensure as a pharmacist; or

(2) A graduate of an approved college of pharmacy or a graduate who has established educational equivalency by obtaining a Foreign Pharmacy Graduate Examination Committee (FPGEC) certificate, who is currently licensed by the board for the purpose of obtaining practical experience as a requirement for licensure as a pharmacist; or
(3) A qualified applicant awaiting examination for licensure; or

(4) An individual participating in a residency or fellowship program.

(l) "Labeling" means the process of preparing and affixing a label to a drug container exclusive, however, of a labeling by a manufacturer, packer or distributor of a nonprescription drug or commercially packaged legend drug or device. Any such label shall include all information required by federal law or regulation and state law or rule.

(m) "Mail order pharmacy" means a pharmacy, regardless of its location, which dispenses greater than ten percent prescription drugs via the mail.

(n) "Manufacturer" means a person engaged in the manufacture of drugs or devices.

(o) "Manufacturing" means the production, preparation, propagation or processing of a drug or device, either directly or indirectly, by extraction from substances of natural origin or independently by means of chemical or biological synthesis and includes any packaging or repackaging of the substance(s) or labeling or relabeling of its contents and the promotion and marketing of such drugs or devices. Manufacturing also includes the preparation and promotion of commercially available products from bulk compounds for resale by pharmacies, practitioners or other persons.

(p) "Nonprescription drug" means a drug which may be sold without a prescription and which is labeled for use by the consumer in accordance with the requirements of the laws and rules of this state and the federal government.

(q) "Patient counseling" means the oral communication by the pharmacist of information, as defined in the rules of the board, to the patient, to improve therapy by aiding in the proper use of drugs and devices.

(r) "Person" means an individual, corporation, partnership, association or any other legal entity, including gov-
(s) "Pharmaceutical care" is the provision of drug therapy and other pharmaceutical patient care services intended to achieve outcomes related to the cure or prevention of a disease, elimination or reduction of a patient's symptoms or arresting or slowing of a disease process as defined in the rules of the board.

(t) "Pharmacist" or "registered pharmacist" means an individual currently licensed by this state to engage in the practice of pharmacy and pharmaceutical care.

(u) "Pharmacist-in-charge" means a pharmacist currently licensed in this state who accepts responsibility for the operation of a pharmacy in conformance with all laws and rules pertinent to the practice of pharmacy and the distribution of drugs and who is personally in full and actual charge of such pharmacy and personnel.

(v) "Pharmacy" means any drugstore, apothecary or place within this state where drugs are dispensed and sold at retail or displayed for sale at retail and pharmaceutical care is provided; and any place outside of this state where drugs are dispensed and pharmaceutical care is provided to residents of this state.

(w) "Pharmacy technician" means registered supportive personnel who work under the direct supervision of a pharmacist who have passed an approved training program as described in this article.

(x) "Practitioner" means an individual currently licensed, registered or otherwise authorized by the jurisdiction in which he or she practices to prescribe and administer drugs in the course of professional practices, including allopathic and osteopathic physicians, dentists, physician's assistants, optometrists, veterinarians, podiatrists and nurse practitioners as allowed by law.

(y) "Preceptor" means an individual who is currently licensed as a pharmacist by the board, meets the qualifications as a preceptor under the rules of the board, and participates in the instructional training of pharmacy interns.
(z) "Prescription drug" or "legend drug" means a drug which, under federal law, is required, prior to being dispensed or delivered, to be labeled with either of the following statements:

1. "Caution: Federal law prohibits dispensing without prescription";
2. "Caution: Federal law restricts this drug to use by, or on the order of, a licensed veterinarian"; or a drug which is required by any applicable federal or state law or rule to be dispensed pursuant only to a prescription drug order or is restricted to use by practitioners only.

(aa) "Prescription drug order" means a lawful order of a practitioner for a drug or device for a specific patient.

(bb) "Prospective drug use review" means a review of the patient's drug therapy and prescription drug order, as defined in the rules of the board, prior to dispensing the drug as part of a drug regimen review.

(cc) "USP-DI" means the United States Pharmacopedia-Dispensing Information.

(dd) "Wholesale distributor" means any person engaged in wholesale distribution of drugs, including, but not limited to, manufacturers' and distributors' warehouses, chain drug warehouses and wholesale drug warehouses; independent wholesale drug trader; and retail pharmacies that conduct wholesale distributions.

§30-5-2. Board of pharmacy; appointment, qualifications and terms of members; compensation; powers and duties generally; meetings and notices.

(a) There shall be a state board of pharmacy, known as the "West Virginia board of pharmacy," which shall consist of five practicing pharmacists and two public members, who shall be appointed by the governor by and with the advice and consent of the Senate. Any vacancy which occurs in the membership of the board for any reason, including expiration of term, removal, resignation, death, disability or disqualification shall be immediately filled by the governor as provided by this section. Nothing in this
section shall require the governor to change the composition of the board prior to the usual expiration of any member's term. The governor may consider the diversity of pharmacy areas of practice when filling vacancies.

(b) Each pharmacist member of the board, at the time of his appointment, shall be a resident of this state, licensed and in good standing to engage in the practice of pharmacy in this state for a period of at least five years prior to their appointment. The public members shall be residents of this state who have attained the age of eighteen years and may not be a past or present pharmacist, the spouse of a pharmacist, a person who has ever had any material financial interest in providing pharmacy services or who has engaged in any activity directly related to the practice of pharmacy.

(c) Each member of the board shall receive two hundred dollars for each day spent in attending to the duties of the board or of its committees, and shall be reimbursed for all actual and necessary expenses incurred in carrying out his or her duties.

(d) The members of the board in office on the date this section takes effect shall, unless sooner removed, continue to serve until their respective terms expire and until their successors have been appointed and have qualified. Board member terms shall be for five years with at least one pharmacist member's term expiring yearly. The governor may, with the advice and consent of the Senate, reappoint any member for additional consecutive terms. Members as of the first day of July, one thousand nine hundred ninety-five, are eligible for reappointment to additional terms regardless of the length of time they have previously served on the board.

(e) The board, in addition to the authority, powers and duties granted to the board by this chapter and chapter sixteen of this code, shall have the authority to:

(1) Regulate the practice of pharmacy;

(2) Regulate the employment of licensed interns in pharmacy;
(3) Appoint, within the limit of appropriations, inspectors who shall be pharmacists, and investigators, to act as agents of the board within the provisions of this chapter and chapter sixteen of this code and rules as the board shall promulgate;

(4) Adopt rules of professional conduct; and

(5) Hire an attorney, as may be necessary.

(f) A majority of the membership of the board constitutes a quorum for the transaction of business, and any motion is approved by a majority vote of a quorum. All board members shall be given advance notice of each board meeting.

(g) Meetings of the board shall be held in public session, except that the board may hold closed sessions to prepare, approve, grade or administer examinations. Disciplinary proceedings, prior to a finding of probable cause, as provided in section seven of this article shall be held in closed sessions, unless the party subject to discipline requests that the hearing be held in public sessions. All discussions or meetings of the board concerning personnel matters shall be held in closed session.

§30-5-2a. Records of board; expungement; examination notice; public information.

(a) The board shall maintain a permanent record of the names of all pharmacists, interns and pharmacy technicians lawfully practicing in this state, and of all persons applying for licensure to practice, along with an individual historical record for each such individual containing reports and all other information furnished to the board concerning any applicant, pharmacist, intern or pharmacy technician.

(b) Upon a determination by the board that any information submitted to it is without merit, the report shall be expunged from the individual's historical record.

(c) Any licensee or registrant of the board or authorized representative thereof, has the right, upon request, to examine his or her own individual historical record main-
tained by the board pursuant to this article and to place
into such record a statement regarding the correctness or
relevance of any information in the historical record.
These statements shall at all times be appended to and
accompany any request for review or copies made of the
portion of the record to which they refer.

d) Orders of the board relating to disciplinary action
against a pharmacist, pharmacy technician, or other license
or registrant of the board are public information.

§30-5-3. When licensed pharmacist required; person not li-
censed pharmacist, pharmacy technician or li-
censed intern not to compound prescriptions or
dispense poisons or narcotics; licensure of in-
terns.

(a) It is unlawful for any person not a pharmacist, or
who does not employ a pharmacist to conduct any phar-
macy, or store for the purpose of retailing, compounding
or dispensing prescription drugs or prescription devices.

(b) It is unlawful for the proprietor of any store or
pharmacy to permit any person not a pharmacist to com-
pound or dispense prescriptions or prescription refills, or
to retail or dispense the poisons and narcotic drugs named
in sections two, three and six, article eight, chapter sixteen
of this code: Provided, That a licensed intern may com-
pound and dispense prescriptions or prescription refills
under the direct supervision of a pharmacist: Provided,
however, That registered pharmacy technicians may assist
in the preparation and dispensing of prescriptions or pre-
scription refills including, but not limited to, reconstitution
of liquid medications, typing and affixing labels under the
direct supervision of a licensed pharmacist.

(c) It is the duty of a pharmacist or employer who
employs an intern, to license the intern with the board
within ninety days after employment. The board shall
furnish proper forms for this purpose and shall issue a
certificate to the intern upon licensure.

(d) The experience requirement for licensure as a
pharmacist shall be computed from the date certified by
the supervising pharmacist as the date of entering the
internship. If the internship is not registered with the
board of pharmacy, then the intern shall receive no credit
for such experience when he or she makes application for
examination for licensure as a pharmacist: Provided, That
credit may be given for such unregistered experience if an
appeal is made and evidence produced showing experi-
ence was obtained but not registered and that failure to
register the internship experience was not the fault of the
intern.

(e) An intern having served part or all of his or her
internship in a pharmacy in another state or foreign coun-
try shall be given credit for the same when the affidavit of
his or her internship is signed by the pharmacist under
whom he or she served, and it shows the dates and number
of hours served in the internship and when the affidavit is
attested by the secretary of the state board of pharmacy of
the state or country where the internship was served.

(f) Up to one third of the experience requirement for
licensure as a pharmacist may be fulfilled by an internship
in a foreign country.

§30-5-4. Use of titles or terms; penalties and fines.

(a) It is unlawful for any person not legally licensed as
a pharmacist, unless he or she employs a licensed pharma-
cist, to take, use or exhibit the title of pharmacist, or li-
censed or registered pharmacist, or the title of druggist or
apothecary, or any other title or description of like import,
or to label, mark, or advertise his or her or any other place
of business as a pharmacy or drugstore or by the use of
the words drug or medicines or any other compound or
derivative of the same, or by any other word or sign indi-
cating or intended to indicate that drugs or pharmaceutical
supplies are either sold or offered for sale.

(b) It is unlawful for any person not legally registered
as a pharmacy technician to take, use or exhibit the title of
pharmacy technician, or any title or description of like
import.

(c) Any person violating this section shall, upon con-
vicition, be deemed guilty of a misdemeanor and fined not
less than five hundred nor more than one thousand dol-

§30-5-5. Qualifications for licensure as pharmacist; fees; cer-
tificates of licensure; rules for licensure; reciproci-

(a) In order to be licensed as a pharmacist within the
meaning of this article, a person shall:

(1) Be eighteen years of age or older;

(2) Present to the board satisfactory evidence that he
or she is a graduate of a recognized school of pharmacy
as defined by the board of pharmacy.

(3) Present to the board satisfactory evidence that he
or she has completed at least fifteen hundred hours of
internship in a pharmacy under the instruction and super-

(4) Pass an examination approved by the board of
pharmacy; and

(5) Present to the board satisfactory evidence that he
or she is a person of good moral character, has not been
convicted of a felony involving controlled substances or
violent crime, and is not addicted to alcohol or the use of
controlled substances.

(b) An applicant for examination shall pay to the
board a fee of one hundred twenty-five dollars with his or
her application.

(c) The board shall issue certificates of licensure to all
persons who successfully pass the required examination
and are otherwise qualified and to all those whose certifi-
cates or licenses the board shall accept in lieu of an exami-
nation as provided in section six of this article.

(d) The board shall by rule stipulate the forms to be
used for licensure application, the requirements for reci-

minimum standards.


§30-5-5a. Legislative findings; registration of pharmacy technicians; qualifications; training programs; rules and restrictions.

(a) The Legislature finds that it is in the best interests of the public health, safety and welfare that licensed pharmacists in this state be assisted with or relieved of certain tasks so that the pharmacist may counsel patients, improve pharmaceutical care and therapeutic outcomes. To achieve this aim, the board shall recognize and register pharmacy technicians.

(b) On or after the first day of July, one thousand nine hundred ninety-six, any person practicing as a pharmacy technician in this state shall be registered with the board of pharmacy pursuant to the provisions of this section.

(c) In order to become registered as pharmacy technicians in this state, individuals shall:

(1) Be at least eighteen years old;

(2) Be a high school graduate or its equivalent;

(3) Present to the board satisfactory evidence that he or she is of good moral character, is not addicted to alcohol or controlled substances and is free of any felony convictions; and

(4) Satisfactorily complete a board-approved pharmacy technician training program.

(d) The pharmacy technician training program and its curriculum shall be designed to train individuals to perform nonprofessional functions as described in legislative rules promulgated in accordance with the provisions of article three, chapter twenty-nine-a of this code.

(e) Pharmacy technicians shall be identified by a name tag and designation as pharmacy technician while working in a pharmacy within this state. A ratio of no more than four pharmacy technicians per on-duty pharmacist operating in any outpatient, mail order or institutional pharmacy shall be maintained.
§30-5-6. Reciprocal licensure of pharmacists from other states or countries.

(a) The board of pharmacy may by reciprocity license pharmacists in this state persons who have been legally registered or licensed pharmacists in another state: Provided, That the applicant for such licensure shall meet the requirements of the rules for reciprocity promulgated by the board in accordance with the provisions of chapter twenty-nine-a of this code: Provided, however, That reciprocity is not authorized for pharmacists from another state where that state does not permit reciprocity to pharmacists licensed in West Virginia.

(b) The board may refuse reciprocity to pharmacists from another country unless the applicant qualifies under such rules as may be promulgated by the board for licensure of foreign applicants.

(c) Applicants for licensure under this section shall, with their application, forward to the secretary of the board of pharmacy a fee of two hundred fifty dollars. In the event the applicant desires to be examined other than at a regular meeting of the board the applicant shall submit to the board an additional fee of one hundred fifty dollars.

§30-5-7. Grounds for suspension or revocation of license or disciplinary proceedings; penalties and procedures; temporary suspensions; reporting of disciplinary action.

(a) The board shall have the power to withhold, revoke or suspend any license or any certificate issued under this article or to penalize or discipline any pharmacist or pharmacy after giving reasonable notice and an opportunity to be heard pursuant to the provisions of section one, article five, chapter twenty-nine-a of this code, any person who has:

1. Become unfit or incompetent to practice pharmacy by reason of: (A) alcohol or substance abuse; (B) insanity; or (C) any abnormal physical or mental condition which threatens the safety of persons to whom such person might
sell or dispense prescriptions, drugs, or devices, or for whom he might manufacture, prepare or package, or supervise the manufacturing, preparation, or packaging of prescriptions, drugs or devices;

(2) Been convicted in any of the courts of this state, the United States of America, or any other state, of a felony or any crime involving moral turpitude which bears a rational nexus to the individual's ability to practice as a pharmacist or pharmacist technician;

(3) Violated any of the provisions of this chapter or chapter sixteen of the code;

(4) Failed to comply with the rules of professional conduct adopted by the board pursuant to section two of this article;

(5) Knowledge or suspicion that a pharmacist, pharmacy technician or pharmacy intern is incapable of engaging in the practice of pharmacy with reasonable skill, competence and safety and has failed to report this information to the board;

(6) Committed fraud as a licensee in connection with the practice of pharmacy;

(7) Performed an act outside this state which would constitute a violation within this state; or

(8) Agreed to participate in a legend drug product conversion program promoted or offered by a manufacturer, wholesaler or distributor of such product for which the pharmacist or pharmacy received any form of financial remuneration, or agreed to participate in a legend drug program in which the pharmacist or pharmacy is promoted or offered as the exclusive provider of legend drug products or whereby in any way the public is denied, limited or influenced in selecting pharmaceutical service or counseling.

(b) Upon a finding of a violation of one or more of the above grounds for discipline by a pharmacist, intern or pharmacy technician, the board may impose one or more of the following penalties:
(1) Suspension of the offender's license or registration for a term to be determined by the board;

(2) Revocation of the offender's license or registration;

(3) Restriction of the offender's license or registration to prohibit the offender from performing certain acts or from engaging in the practice of pharmacy in a particular manner for a term to be determined by the board;

(4) Imposition of a fine not to exceed one thousand dollars for each offense;

(5) Refusal to renew the offender's license or registration;

(6) Placement of the offender on probation and supervision by the board for a period to be determined by the board.

(c) All final decisions of the board shall be subject to judicial review pursuant to the procedures of article five, chapter twenty-nine-a of this code.

(d) In the case of a pharmacy or wholesale distributor, the disciplinary order may be entered as to the corporate owner, if any, as well as to the pharmacist, officer, owner or partner of the pharmacy or wholesale distributor if it is found that such person or entity had knowledge of or knowingly participated in one or more of the violations set forth in this article or of article three, chapter sixty-a of this code.

(e) Notwithstanding the provisions of section eight, article one, chapter thirty of this code, if the board determines that the evidence in its possession indicates that a pharmacist's continuation in practice or unrestricted practice constitutes an immediate danger to the public, the board may, on a temporary basis and without a hearing, take any of the actions provided for in this section if proceedings for a hearing before the board are initiated simultaneously with the temporary action and begin within fifteen days of such action. The board shall render its decision within five days of the conclusion of a hearing conducted pursuant to the provisions of this section.
(f) In every disciplinary or licensure case considered by the board pursuant to this article, whether initiated by the board or upon complaint or information from any person or organization, the board shall make a preliminary determination as to whether probable cause exists to substantiate charges of disqualification due to any reason set forth in this section. If such probable cause is found to exist, all proceedings on such charges shall be open to the public, who shall be entitled to all reports, records and nondeliberative materials introduced at such hearing, including the record of any final action taken: Provided, That any medical records pertaining to a person who has not expressly waived his or her right to the confidentiality of such records shall not be open to the public.

(g) All disciplinary actions taken by the board shall be reported to the national board of pharmacy, appropriate federal agencies and to any other state boards with which the disciplined licensee may also be registered or licensed.

§30-5-7a. Required reporting of information to board pertaining to professional malpractice and convictions; complaints of professional incompetence; reporting forms.

(a) Every person, partnership, corporation, association, insurance company, professional society or other organization providing professional liability insurance to a pharmacist, pharmacist technician or intern in this state shall submit to the board the following information within thirty days from any judgment, dismissal or settlement of a civil action or of any claim involving the insured: The date of any judgment or settlement; the amount of any settlement or judgment against the insured; and such other information as the board may require.

(b) Within thirty days after a person known to be a pharmacist, pharmacy intern, or pharmacy technician licensed or otherwise lawfully practicing pharmacy in this state or applying to be so licensed is convicted of any crime under the laws of this state, or the laws of the United States which involves drugs in any way, including any controlled substance under state or federal law, the clerk of the court of record in which the conviction was entered
shall forward to the board a certified true and correct
abstract of record of the convicting court. The abstract
shall include the name and address of such licensee, the
nature of the offense committed and the final judgment
and sentence of the court.

(c) Any person may report to the board relevant facts
about the conduct of a licensee of the board which in the
opinion of such person amounts to professional malprac-
tice or professional incompetence.

(d) The board shall provide forms for filing reports
pursuant to this section. Reports submitted in other forms
shall be accepted by the board.

§30-5-7b. Voluntary agreements relating to alcohol or chemical dependency; confidentiality of same.

(a) In order to encourage voluntary reporting of alco-
hol or other chemical dependency impairment and in
recognition of the fact that alcoholism and chemical de-
pendency are illnesses, a pharmacist or pharmacy techni-
cian or other licensee or registrant or the board may enter
into a voluntary agreement with the board reporting his or
her participation in an alcohol or chemical dependency
treatment program or reporting an alcohol or chemical
dependency impairment to the board and seek treatment
for his or her dependency. Pursuant to said agreement, the
board shall impose limitations on the practice of said
pharmacist, pharmacy technician or other licensee or reg-
istrant of the board.

(b) Any voluntary agreement entered into pursuant to
this subsection may not be considered a disciplinary ac-
tion or order by the board and shall not be public infor-
mation if:

(1) Such voluntary agreement is the result of the phar-
macist, pharmacy technician, or other licensee or registrant
of the board reporting his or her participation in an alco-
hol or chemical dependency treatment program or report-
ing to the board his or her alcohol or chemical dependen-
cy impairment and requesting such an agreement for the
purpose of seeking treatment; and
(2) The board has not received nor filed any written complaints regarding said pharmacist, pharmacy technician or other licensee or registrant of the board relating to an alcohol or chemical dependency impairment affecting the care and treatment of patients or customers, nor received any reports pursuant to section seven of this article relating to an alcohol or chemical dependency impairment.

(c) If any pharmacist, pharmacy technician or other licensee or registrant enters into a voluntary agreement with the board pursuant to this subsection and then fails to comply with or fulfill the terms of said agreement, the board shall initiate disciplinary proceedings pursuant to section seven of this article.

(d) If the board has not instituted any disciplinary proceedings as provided for in this article, any information received, maintained or developed by the board relating to the alcohol or chemical dependency impairment of any pharmacist or pharmacy technician, other licensee or registrant of the board and any voluntary agreement made pursuant to this subsection shall be confidential and not available for public information, discovery or court subpoena nor for introduction into evidence in any professional liability action or other action for damages arising out of the provision of or failure to provide health care services.

(e) In the board's annual report of its activities to the Legislature required under section eight of this article, the board shall include information regarding the success of the voluntary agreement mechanism established therein: Provided, That in making such report the board shall not disclose any personally identifiable information relating to any pharmacist or other licensee or registrant of the board participating in a voluntary agreement as provided herein.

(f) Notwithstanding any of the foregoing provisions, the board may cooperate with and provide documentation of any voluntary agreement entered into pursuant to this subsection to licensing boards in other jurisdictions, as may be appropriate.
(g) Any restrictions on the disclosure of confidential information does not apply to any investigation or proceeding by the board or by a hospital governing board or committee with respect to relevant medical records, while any of the aforesaid are acting within the scope of their authority as stated in law or in the hospital bylaws, rules, regulations or policies and procedures: Provided, That the disclosure of any information pursuant to this provision shall not be considered a waiver of any such privilege in any other proceeding.

§30-5-8. Reports by secretary of board to secretary of state; "list of pharmacists."

The secretary of the board of pharmacy shall provide the secretary of state with a list of all pharmacists, pharmacy technicians and pharmacy interns in this state, giving the name of the person, his or her business address, and the date of his or her licensure registration. On or before the fifteenth day of September each year, the secretary of the board shall certify to the secretary of state all changes in said list required by the addition of new licensures, registrations, renewals, reported deaths, forfeitures of licenses or registrations or for other causes, occurring during the preceding year. The secretary of state shall enter in an appropriate book, known as "List of Pharmacists" the facts shown by such reports, which reports shall be filed and preserved in his or her office.

§30-5-9. Fees.

The board of pharmacy shall charge and collect the following fees, in addition to those provided in article one of this chapter and in sections five, fourteen and sixteen of this article: For renewing the licensure of a pharmacist, thirty dollars; to license an intern pharmacist, ten dollars plus five dollars for each of the remaining periods of his or her internship; to register a consultant pharmacist, twenty dollars for the initial application and ten dollars for each additional application; and to register a pharmacy technician, twenty-five dollars and ten dollars for each renewal.

§30-5-10. Annual renewal of license; fees and notices.
(a) Every licensed pharmacist, intern or pharmacy technician who desires to renew his or her license shall on or before the first day of July, one thousand nine hundred ninety-one, and annually thereafter apply to the state board of pharmacy for a renewal of his or her license, and shall transmit with his or her application the fee prescribed in the preceding section of this article. Notification of the annual renewal shall be given by the board at least thirty days prior to said first day of July. Such notification shall be mailed to the last known address of each pharmacist or pharmacy technician as shown on record with the board.

(b) If any pharmacist or pharmacy technician fails for a period of sixty days after the first day of July of each year to apply to the board for a renewal of his or her license, the board shall send a second notification of the required annual renewal to the last known address of the pharmacist or pharmacy technician by certified mail, return receipt requested. If the pharmacist or pharmacy technician fails to apply to the board for a renewal of his or her license within thirty days after receipt of the second notification, his or her name shall be erased from the register of pharmacists and pharmacy technicians.

(c) In order for any pharmacist or pharmacy technician whose name has been erased from the register of the board pursuant to subsection (b) of this section to again become licensed, such pharmacist or pharmacy technician shall appear personally before the board, or an authorized committee of the board, to show cause for permitting the license to lapse. If such person submits to the board satisfactory reasons for allowing the license to lapse and satisfies the board as to his or her qualifications to practice the profession, such person shall be reinstated upon payment of a reinstatement fee of two hundred fifty dollars plus the renewal fee of thirty dollars.

§30-5-11. Certificate of licensure or permit shall be displayed.

Every certificate of registration or licensure to practice as a pharmacist, intern or pharmacy technician, and every renewal of such certificate or permit, shall be conspicuously displayed in the pharmacy or place of business of which the pharmacist, intern or pharmacy technician or
other person to whom it is issued is the owner or manager, or in which he or she is employed.

§30-5-12. Responsibility for quality of drugs dispensed; exception; falsification of labels; deviation from prescription.

(a) All persons, whether licensed pharmacists or not, shall be responsible for the quality of all drugs, chemicals and medicines they may sell or dispense, with the exception of those sold in or dispensed unchanged from the original retail package of the manufacturer, in which event the manufacturer shall be responsible.

(b) Except as provided in section twelve-b of this article, the following acts shall be prohibited: (1) The falsification of any label upon the immediate container, box and/or package containing a drug; (2) the substitution or the dispensing of a different drug in lieu of any drug prescribed in a prescription without the approval of the practitioner authorizing the original prescription: Provided, That this shall not be construed to interfere with the art of prescription compounding which does not alter the therapeutic properties of the prescription or appropriate generic substitute; (3) the filling or refilling of any prescription for a greater quantity of any drug or drug product than that prescribed in the original prescription without a written order or an oral order reduced to writing, or the refilling of a prescription without the verbal or written consent of the practitioner authorizing the original prescription.

§30-5-12b. Definitions; selection of generic drug products; exceptions; records; labels; manufacturing standards; rules; notice of substitution; complaints; notice and hearing; immunity.

(a) As used in this section:

(1) "Brand name" means the proprietary or trade name selected by the manufacturer and placed upon a drug or drug product, its container, label or wrapping at the time of packaging.

(2) "Generic name" means the official title of a drug
or drug combination for which a new drug application, or
an abbreviated new drug application, has been approved
by the United States food and drug administration and is
in effect.

(3) "Substitute" means to dispense without the
prescriber's express authorization a therapeutically equiva-
ient generic drug product in the place of the drug ordered
or prescribed.

(4) "Equivalent" means drugs or drug products which
are the same amounts of identical active ingredients and
same dosage form, and which will provide the same thera-
peutic efficacy and toxicity when administered to an indi-
vidual and is approved by the United States food and drug
administration.

(5) "Practitioner" means a physician, an authorized
Type A physician assistant at the direction of his or her
supervising physician in accordance with the provisions of
section sixteen, article three of this chapter, osteopath,
dentist, veterinarian, podiatrist, optometrist or any other
person duly licensed to practice and to prescribe drugs
under the laws of this state.

(b) A pharmacist who receives a prescription for a
brand name drug or drug product shall substitute a less
expensive equivalent generic name drug or drug product
unless in the exercise of his or her professional judgment
the pharmacist believes that the less expensive drug is not
suitable for the particular patient: Provided, That no sub-
stitution may be made by the pharmacist where the pre-
scribing practitioner indicates that, in his or her profes-
sonal judgment, a specific brand name drug is medically
necessary for a particular patient.

(c) A written prescription order shall permit the phar-
macist to substitute an equivalent generic name drug or
drug product except where the prescribing practitioner has
indicated in his or her own handwriting the words "Brand
Medically Necessary." The following sentence shall be
printed on the prescription form: "This prescription may
be filled with a generically equivalent drug product unless
the words 'Brand Medically Necessary' are written, in the
practitioner's own handwriting, on this prescription form.":
Provided, That "Brand Medically Necessary" may be indi-
cated on the prescription order other than in the prescribing practitioner's own handwriting unless otherwise required by federal mandate.

(d) A verbal prescription order shall permit the pharmacist to substitute an equivalent generic name drug or drug product except where the prescribing practitioner shall indicate to the pharmacist that the prescription is "Brand Necessary" or "Brand Medically Necessary." The pharmacist shall note the instructions on the file copy of the prescription or chart order form.

(e) No person may by trade rule, work rule, contract, or in any other way prohibit, restrict, limit or attempt to prohibit, restrict or limit the making of a generic name substitution under the provisions of this section. No employer or his or her agent may use coercion or other means to interfere with the professional judgment of the pharmacist in deciding which generic name drugs or drug products shall be stocked or substituted: Provided, That this section shall not be construed to permit the pharmacist to generally refuse to substitute less expensive therapeutically equivalent generic drugs for brand name drugs, and that any pharmacist so refusing shall be subject to the penalties prescribed in section twenty-two, article five, chapter thirty of this code.

(f) A pharmacist may substitute a drug pursuant to the provisions of this section only where there will be a savings to the buyer. Where substitution is proper pursuant to this section, or where the practitioner prescribes the drug by generic name, the pharmacist shall, consistent with his or her professional judgment, dispense the lowest retail cost, effective brand which is in stock.

(g) All savings in the retail price of the prescription shall be passed on to the purchaser; these savings shall be equal to the difference between the retail price of the brand name product and the customary and usual price of the generic product substituted therefor: Provided, That in no event shall such savings be less than the difference in acquisition cost of the brand name product prescribed and the acquisition cost of the substituted product.
(h) Each pharmacy shall maintain a record of any substitution of an equivalent generic name drug product for a prescribed brand name drug product on the file copy of a written or verbal prescription or chart order. Such record shall include the manufacturer and generic name of the drug product selected.

(i) All drugs shall be labeled in accordance with the instructions of the practitioner.

(j) Unless the practitioner directs otherwise, the prescription label on all drugs dispensed by the pharmacist shall indicate the generic name using abbreviations if necessary and either the name of the manufacturer or packager, whichever is applicable in the pharmacist's discretion. The same notation will be made on the original prescription retained by the pharmacist.

(k) A pharmacist may not dispense a product under the provisions of this section unless the manufacturer has shown that the drug has been manufactured with the following minimum good manufacturing standards and practices by:

   (1) Labeling products with the name of the original manufacturer and control number;

   (2) Maintaining quality control standards equal to or greater than those of the United States Food and Drug Administration;

   (3) Marking products with identification code or monogram; and

   (4) Labeling products with an expiration date.

(l) The West Virginia board of pharmacy shall promulgate rules in accordance with the provisions of chapter twenty-nine-a of this code which establish a formulary of generic type and brand name drug products which are determined by the board to demonstrate significant biological or therapeutic inequivalence and which, if substituted, would pose a threat to the health and safety of patients receiving prescription medication. The formulary shall be promulgated by the board within ninety days of
the date of passage of this section, and may be amended in
accordance with the provisions of chapter twenty-nine-a of
this code.

(m) No pharmacist shall substitute a generic named
therapeutically equivalent drug product for a prescribed
brand name drug product if the brand name drug product
or the generic drug type is listed on the formulary estab-
lished by the West Virginia board of pharmacy pursuant
to this article, or is found to be in violation of the require-
ments of the United States Food and Drug Administration.

(n) Any pharmacist who substitutes any drug shall,
either personally or through his or her agent, assistant or
employee, notify the person presenting the prescription of
such substitution. The person presenting the prescription
shall have the right to refuse the substitution. Upon re-
quest the pharmacist shall relate the retail price difference
between the brand name and the drug substituted for it.

(o) Every pharmacy shall post in a prominent place
that is in clear and unobstructed public view, at or near the
place where prescriptions are dispensed, a sign which shall
read: "West Virginia law requires pharmacists to substitute
a less expensive generic named therapeutically equivalent
drug for a brand name drug, if available, unless you or
your physician direct otherwise." The sign shall be printed
with lettering of at least one and one-half inches in height
with appropriate margins and spacing as prescribed by the
West Virginia board of pharmacy.

(p) The West Virginia board of pharmacy shall pro-
mulgate rules in accordance with the provisions of chapter
twenty-nine-a of this code setting standards for substituted
drug products, obtaining compliance with the provisions
of this section and enforcing the provisions of this section.

(q) Any person shall have the right to file a complaint
with the West Virginia board of pharmacy regarding any
violation of the provisions of this article. Such complaints
shall be investigated by the board of pharmacy.

(r) Fifteen days after the board has notified, by regis-
tered mail, a person, firm, corporation or copartnership
that such person, firm, corporation or copartnership is suspected of being in violation of a provision of this section, the board shall hold a hearing on the matter. If, as a result of the hearing, the board determines that a person, firm, corporation or copartnership is violating any of the provisions of this section, it may, in addition to any penalties prescribed by section twenty-two of this article, suspend or revoke the permit of any person, firm, corporation or copartnership to operate a pharmacy.

(s) No pharmacist complying with the provisions of this section shall be liable in any way for the dispensing of a generic named therapeutically equivalent drug, substituted under the provisions of this section, unless the generic named therapeutically equivalent drug was incorrectly substituted.

(t) In no event where the pharmacist substitutes a drug under the provisions of this section shall the prescribing physician be liable in any action for loss, damage, injury or death of any person occasioned by or arising from the use of the substitute drug unless the original drug was incorrectly prescribed.

(u) Failure of a practitioner to specify that a specific brand name is necessary for a particular patient shall not constitute evidence of negligence unless the practitioner had reasonable cause to believe that the health of the patient required the use of a certain product and no other.

§30-5-13. Each pharmacy to have USP-DI.

Every pharmacy as defined in this article, shall own and have in the pharmacy at all times in text or electronic form, a recent edition of the USP-DI and any supplements. No license or renewal shall be issued until a USP-DI is in the pharmacy.

§30-5-14. Pharmacies to be registered; permit to operate; fees; pharmacist to conduct business.

(a) The board of pharmacy shall require and provide for the annual registration of every pharmacy doing business in this state. Any person, firm, corporation or partnership desiring to operate, maintain, open or establish a
pharmacy in this state shall apply to the board of pharmacy for a permit to do so. The application for such permit shall be made on a form prescribed and furnished by the board of pharmacy, which, when properly executed, shall indicate the owner, manager, trustee, lessee, receiver, or other person or persons desiring such permit, as well as the location of such pharmacy, including street and number, and such other information as the board of pharmacy may require. If it is desired to operate, maintain, open or establish more than one pharmacy, separate application shall be made and separate permits or licenses shall be issued for each.

(b) Every initial application for a permit shall be accompanied by the required fee of one hundred fifty dollars. The fee for renewal of such permit or license shall be seventy-five dollars annually.

(c) If an application is approved, the secretary of the board of pharmacy shall issue to the applicant a permit or license for each pharmacy for which application is made. Permits or licenses issued under this section shall not be transferable and shall expire on the thirtieth day of June of each calendar year, and if application for renewal of permit or license is not made on or before that date, or a new one granted on or before the first day of August, following, the old permit or license shall lapse and become null and void and shall require an inspection of the pharmacy and a fee of one hundred fifty dollars plus one hundred fifty dollars for the inspection.

(d) Every such place of business so registered shall employ a pharmacist in charge and operate in compliance with the general provisions governing the practice of pharmacy and the operation of a pharmacy.

(e) The provisions of this section shall have no application to the sale of nonprescriptive drugs which are not required to be dispensed pursuant to a practitioner's prescription.


(a) Every pharmacy at all times, shall be under the
direction and supervision of a licensed pharmacist who shall be designated by the owner of the pharmacy as the pharmacist-in-charge. This designation must be filed with the board within thirty days of the designation.

(b) The pharmacist-in-charge is responsible for the pharmacy's compliance with state and federal pharmacy laws and regulations and for maintaining records and inventory.

(c) It is a violation of this section if the owner of a pharmacy fails to designate a pharmacist-in-charge or permits the practice of pharmacy without having designated a pharmacist-in-charge, or fails to notify the board of pharmacy if the designated pharmacist-in-charge leaves the employ of the pharmacy.

(d) Before a permit is issued to operate a pharmacy, or renewed, the application shall designate the pharmacist-in-charge. The designated pharmacist-in-charge shall be present when a new store is to be inspected.

(e) A pharmacist-in-charge shall not hold such designated position at more than one pharmacy, whether within or without the state of West Virginia. The board of pharmacy shall promulgate rules in accordance with the provisions of chapter twenty-nine-a of this code relative to pharmacies which are operated over forty hours a week.

(f) An interim pharmacist-in-charge may be designated for a period not to exceed sixty days. The request for an interim pharmacist-in-charge shall detail the circumstances which warrant such a change. This change in designation shall be filed with the board within thirty days of the designation.

(g) The board of pharmacy shall furnish the form which designates a change of the pharmacist-in-charge and every such application shall be subject to a fee of ten dollars.

§30-5-15. Professional and technical equipment required for pharmacy or drugstore; penalties and fines.

(a) Every pharmacy shall be equipped with proper
pharmaceutical utensils so that prescriptions can be properly filled and compounded. The board of pharmacy shall by rule prescribe the minimum equipment which a pharmacy shall possess.

(b) Any person violating this section is guilty of a misdemeanor and shall be fined not less than two hundred fifty dollars nor more than one thousand dollars, and no permit shall be issued or renewed for any pharmacy which has not complied with the provisions of this section.

§30-5-16. Permit for manufacture and packaging of drugs, medicines, cosmetics; distribution of legend drugs; regulations as to sanitation and equipment; penalties; revocation of permit.

(a) No drugs or medicines, or toilet articles, dentifrices, or cosmetics, shall be manufactured, made, produced, packed, packaged or prepared within the state, except under the personal supervision of a pharmacist or such other person as may be approved by the board of pharmacy, after an investigation and determination by the board that they are qualified by scientific or technical training and/or experience to perform such duties of supervision as may be necessary to protect the public health and safety.

(b) No person shall manufacture, make, produce, pack, package or prepare any such articles without first obtaining a permit to do so from the board of pharmacy. The permit shall be subject to such rules with respect to sanitation and/or equipment, as the board of pharmacy may from time to time adopt for the protection of the public health and safety.

(c) Any person, firm, corporation, partnership, company, cooperative society or organization who offers for sale, sells, offers or exposes for sale through the method of distribution any legend drugs shall be subject to this article.

(d) The application for any permit required by this section shall be made on a form to be prescribed and furnished by the board of pharmacy and shall be accompanied by the following fees: For a distributor, one hun-
dred fifty dollars, for a manufacturer, five hundred dollars, which amounts shall also be paid as the fees for each annual renewal of such permits. Separate applications shall be made and separate permits issued for each separate place of manufacture, distribution, making, producing, packing, packaging or preparation.

(e) The following fees shall be charged for a permit to handle controlled substances: For a hospital or clinic, fifty dollars; for extended care facilities, twenty-five dollars; for a nursing home, twenty-five dollars; for a teaching institution, twenty-five dollars; for a researcher, twenty-five dollars; for a medical examiner, twenty-five dollars; and for a pharmacy or drug store, fifteen dollars, which amounts shall also be paid for each annual renewal of such permits.

(f) Permits issued under the provisions of this section shall be posted in a conspicuous place in the factory or place for which issued; such permits shall not be transferable, and shall expire on the thirtieth day of June following the day of issue and shall be renewed annually. Nothing in this section shall be construed to apply to those operating registered pharmacies.

(g) Any person, firm, corporation, partnership, company, cooperative society or organization violating any of the provisions of this section and any permittee hereunder who shall violate any of the conditions of this permit or any of the rules adopted by the board of pharmacy shall, upon conviction, be deemed guilty of a misdemeanor and fined not more than fifty dollars for each offense. Each and every day such violation continues shall constitute a separate and distinct offense. Upon conviction of a permittee, his permit shall also immediately be revoked and become null and void.

(h) Any person, firm, corporation, partnership, company, cooperative society, organization or any permittee who is convicted of two or more successive violations of the provisions of this section or of the rules adopted by the board of pharmacy shall at the discretion of the board of pharmacy have such permit permanently revoked, and the board of pharmacy shall refuse to issue further permits to such person, firm, corporation, partnership, company,
§30-5-16b. Partial filling of prescriptions.

The partial filling of a prescription for a controlled substance listed in Schedule II is permissible if the pharmacist is unable to supply the full quantity called for in a written or emergency oral prescription and he makes a notation of the quantity supplied on the face of the written prescription or on the written record of the emergency oral prescription. The remaining portion of the prescription may be filled within seventy-two hours of the first partial filling: Provided, That if the remaining portion is not or cannot be filled within the seventy-two hour period, the pharmacist shall so notify the prescribing individual practitioner. No further quantity may be supplied beyond seventy-two hours without a new prescription.

§30-5-16c. Partial filling of prescriptions for long-term care facility or terminally ill patients; requirements; records; violations.

(a) As used in this section, "long-term care facility" or "LTCF" means any nursing home, personal care home, or residential board and care home as defined in section two, article five-c, chapter sixteen of this code which provides extended health care to resident patients: Provided, That the care or treatment in a household, whether for compensation or not, of any person related by blood or marriage, within the degree of consanguinity of second cousin to the head of the household, or his or her spouse, may not be deemed to constitute a nursing home, personal care home or residential board and care home within the meaning of this article. This section shall not apply to:

(1) Hospitals, as defined under section one, article five-b, chapter sixteen of this code or to extended care facilities operated in conjunction with a hospital;

(2) State institutions as defined in section six, article one, chapter twenty-seven or in section three, article one, chapter twenty-five, all of this code;

(3) Nursing homes operated by the federal government;
(4) Facilities owned or operated by the state government;

(5) Institutions operated for the treatment and care of alcoholic patients;

(6) Offices of physicians; or

(7) Hotels, boarding homes or other similar places that furnish to their guests only a room and board.

(b) As used in this section, "terminally ill" means that an individual has a medical prognosis that his life expectancy is six months or less.

(c) Schedule II prescriptions for patients in a LTCF and for terminally ill patients shall be valid for a period of sixty days from the date of issue unless terminated within a shorter period by the discontinuance of the medication.

(d) A prescription for a Schedule II controlled substance written for a patient in a LTCF or for a terminally ill patient may be filled in partial quantities, including, but not limited to, individual dosage units. The total quantity of Schedule II controlled substances dispensed in all partial filling shall not exceed the total quantity prescribed.

(1) If there is any question whether a patient may be classified as having a terminal illness, the pharmacist shall contact the prescribing practitioner prior to partially filling the prescription.

(2) Both the pharmacist and the prescribing practitioner have a corresponding responsibility to assure that the controlled substance is for a terminally ill patient.

(e) The pharmacist shall record on the prescription that the patient is "terminally ill" or a "LTCF patient". A prescription that is partially filled and does not contain the notation "terminally ill" or "LTCF patient" shall be deemed to have been filled in violation of section three hundred eight, article three, chapter sixty-a of this code.

(f) For each partial filling, the dispensing pharmacist shall record on the back of the prescription, or on another appropriate record which is readily retrievable, the follow-
ing information:

(1) The date of the partial filling;
(2) The quantity dispensed;
(3) The remaining quantity authorized to be dispensed; and
(4) The identification of the dispensing pharmacist.

(g) Information pertaining to current Schedule II prescriptions for terminally ill and LTCF patients may be maintained in a computerized system if such a system has the capability to permit either by display or printout, for each patient and each medication, all of the information required by this section as well as the patient's name and address, the name of each medication, original prescription number, date of issue, and prescribing practitioner information. The system shall also allow immediate updating of the prescription record each time a partial filling of the prescription is performed and immediate retrieval of all information required under this section.

§30-5-19. Rules of board of pharmacy; revocation of permits; employment of field agents, chemists, clerical and other qualified personnel.

(a) The board of pharmacy shall promulgate rules in accordance with the provisions of chapter twenty-nine-a of this code not inconsistent with law, as are necessary to carry out the purposes and enforce the provisions of this article. The board may revoke any permit or license issued under the provisions of this article at any time when examination or inspection of the pharmacy discloses that such place of business is not being conducted according to law.

(b) The board of pharmacy shall have the power and authority to employ field agents, chemists, clerical help, hearing examiners and other qualified personnel, as may be necessary to carry out the purposes and enforce the provisions of this article.

§30-5-21. Limitations of article.

(a) Nothing in this article shall be construed to pre-
vent, restrict or in any manner interfere with the sale of
non-narcotic nonprescription drugs which may be lawful-
ly sold without a prescription in accordance with the Unit-
ed States food, drug, and cosmetic act, or the laws of this
state, nor shall any rule be adopted by the board which
shall require the sale of nonprescription drugs by a li-
censed pharmacist or in a pharmacy, or which shall pre-
vent, restrict, or otherwise interfere with the sale or distri-
bution of such drugs by any retail merchant. The sale or
distribution of nonprescription drugs shall not be deemed
to be improperly engaging in the practice of pharmacy.

(b) Nothing in this article shall be construed to inter-
fere with any legally qualified practitioner of medicine,
dentistry or veterinary medicine, who is not the proprietor
of the store for the dispensing or retailing of drugs, and
who is not in the employ of such proprietor, in the com-
pounding of his own prescriptions, or to prevent him from
supplying to his patients such medicines as he may deem
proper, if such supply is not made as a sale.

§30-5-22. Offenses; penalties.

(a) Any person who violates any of the provisions of
section three of this article is guilty of a misdemeanor,
and, upon conviction, shall, for each offense, be fined not
less than two hundred fifty dollars nor more than one
thousand dollars, or confined in the county jail not to
exceed six months, or both fined and imprisoned, in the
discretion of the court, and each day such violation shall
continue shall be deemed a separate offense.

(b) Any person who violates any of the provisions of
section twelve is guilty of a misdemeanor, and, upon con-
viction, shall be punished by a fine of not less than fifty
nor more than one hundred fifty dollars for each offense.

(c) Any person, except for the board of pharmacy or
board member acting within the scope of his or her re-
sponsibilities or duties as such member, who violates any
of the provisions of section twelve-b is guilty of a misde-
meanor, and, upon conviction, shall be punished by a fine
of not less than fifty nor more than one thousand dollars
for each offense.
(d) Any person, firm, partnership or corporation who violates any of the provisions of section fourteen is guilty of a misdemeanor, and, upon conviction, for the first offense shall be fined not to exceed one hundred dollars, or shall be imprisoned in the county jail not to exceed six months, or both fined and imprisoned, in the discretion of the court. Each and every day that the violation continues shall constitute a separate offense.

(e) Any person, firm, partnership or corporation who violates any of the provisions of section eighteen is guilty of a misdemeanor, and, upon conviction, shall be fined not to exceed fifty dollars for the first offense, and upon conviction of a second offense shall be fined not less than fifty nor more than five hundred dollars, or shall be imprisoned in the county jail not to exceed thirty days, or both fined and imprisoned. Each and every day that the violation continues shall constitute a separate offense.

§30-5-22a. Civil immunity for board members; liability limitations of professionals reporting to board; reporting results of litigation to the board; rules.

(a) The members of the board when acting in good faith and without malice shall enjoy immunity from individual civil liability while acting within the scope of their duties as board members.

(b) Any licensee of this board who reports or otherwise provides evidence of the negligence, impairment or incompetence of another member of this profession to the board or to any peer review organization, shall not be liable to any person for making such a report if such report is made without actual malice and in the reasonable belief that such report is warranted by the facts known to him or her at the time.

(c) Within thirty days of the dismissal, settlement, adjudication or other termination of any claim or cause of action asserted against any professional reporting under the provisions of this article the person or persons filing such claim or cause of actions shall submit to the board the following information:
19   (1) The names of the parties involved;
20   (2) The name of the court in which the action was
21       filed, if applicable;
22   (3) The basis and nature of the claim or cause of ac-
23       tion; and
24   (4) The results of such claim or cause of action, in-
25       cluding dismissal, settlement, court or jury verdict or other
26       means of termination.
27   (e) The board shall promulgate legislative rules in
28       accordance with the provisions of chapter twenty-nine-a of
29       this code establishing procedures for imposing sanctions
30       and penalties against any licensee who fails to submit to
31       the board the information required by this section.
The Joint Committee on Enrolled Bills hereby certifies that the foregoing bill is correctly enrolled.

[Signature]
Chairman Senate Committee

[Signature]
Chairman House Committee

Originating in the House.

Takes effect ninety days from passage.

[Signature]
Clerk of the Senate

[Signature]
Clerk of the House of Delegates

[Signature]
President of the Senate

[Signature]
Speaker of the House of Delegates

The within _______________ this the ____________

day of ____________________, 1995.

[Signature]
Governor