RULES
OF
THE TENNESSEE BOARD OF PHARMACY

CHAPTER 1140-01
INTRODUCTORY RULES

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1140-01-.01 DEFINITIONS.

(1) “ACPE” means the Accreditation Council for Pharmaceutical Education.

(2) “Alternate or alternative infusion pharmacy practice site” means a pharmacy practice site where parenteral, enteral or respiratory therapies, and ancillary supplies, medications and equipment are provided to patients in a non-institutional setting.

(3) “Accreditation Council for Pharmacy Education (ACPE)” means the national organization for accreditation of professional degree programs in pharmacy and for accreditation of providers of continuing pharmacy education.

(4) “Blood” means whole blood collected from a single donor and processed either for transfusion or further manufacturing.

(5) “Blood fraction/component” means that part of blood separated by physical or mechanical means.

(6) “Centralized Prescription Processing” is the filling or refilling of a lawful prescription order written by the patient’s authorized prescriber by one (1) pharmacy licensed by the State of Tennessee at the request of another pharmacy licensed by the State of Tennessee for the delivery of the prescription drugs to the patient or patient’s agent.

(7) “Certified pharmacy technician” means an individual who is certified by a national or state agency that offers a certification program that is recognized by the board.

(8) “Commercially available” means any marketed FDA-approved drug or biologic product not currently listed on any official shortage list recognized by the Board of Pharmacy.

(9) “Component” means any active ingredient, or any added substance, inactive ingredient, excipient or pharmaceutic ingredient, intended for use in the compounding of a drug product including those that may not appear on the product label.

(10) “Consultant pharmacist” means a pharmacist retained on a routine basis to consult with organizations, institutional facilities or patients in areas that pertain to the practice of pharmacy.
“Contact hour” means any hour of completed continuing pharmaceutical education programming which is:

(a) accredited by ACPE (including, but not limited to, live programs, independent study courses, home correspondence courses, and audio or video cassettes); or

(b) approved by the board (including, but not limited to, attendance at state, district, or local pharmacy association meetings).

“Continuing education unit” means ten (10) hours of participation in an ACPE approved or board-approved continuing pharmaceutical education program under responsible sponsorship, capable direction, and qualified instruction.

“Drug sample” means a unit of a prescription drug that is not intended to be sold and is intended to promote the sale of the prescription drug.

“Electronic medical or prescription order” means a medical or prescription order which is transmitted by computer technology other than by electronic image transmission.

“Facsimile (FAX) medical or prescription order” means a medical or prescription order which is transmitted by an electronic image transmission.

“Foreign pharmacy graduate” means a person whose undergraduate pharmacy degree was conferred by any college or school of pharmacy not accredited by the ACPE but which is listed in the World Health Organization World Directory of Colleges and Schools of Pharmacy, or otherwise approved by the Foreign Pharmacy Graduate Examination Committee (FPGEC) certification program as established by the National Association of Boards of Pharmacy.

“Hazardous product” means any substance that may be cytotoxic, genotoxic, oncogenic, mutagenic, teratogenic, or otherwise pose a potential health hazard.

“Institutional facility” means any organization whose primary purpose is to provide a physical environment for patients to obtain health care services, and where patients spend a majority of their time within the facility, including but not limited to a(n):

(a) adult care facility;
(b) assisted living facility;
(c) correctional facility;
(d) developmental disability center;
(e) hospital;
(f) inpatient psychiatric center;
(g) intermediate care facility for the mentally retarded;
(h) mental health facility;
(i) nursing facility;
(j) personal care home;
(k) rehabilitation center;
(l) residential drug or alcohol treatment center;
(m) rest home;
(n) retirement center;
(o) sub-acute care facility; and
(p) university health center.
(Rule 1140-01-.01, continued)

(19) "Institutional pharmacy practice site" means a pharmacy practice site serving patients within an institutional facility.

(20) "Medication order" means a prescription order for any prescription drug or device or related material issued by an authorized prescriber to authorized healthcare personnel in an institutional facility or institutional pharmacy practice site.

(21) "National Association of Boards of Pharmacy (NABP)" means the professional organization that represents the individual state boards of pharmacy.

(22) "Nuclear pharmacy practice site" means a pharmacy practice site providing radiopharmaceutical services.

(23) "Patient counseling" means communication by the pharmacist of information to the patient or caregiver in order to improve therapeutic outcome.

(24) "Pharmaceutical care" is the responsible provision of drug therapy through, among other things, pharmacists identifying potential and actual drug-related problems and resolving and preventing drug-related problems, for the purpose of achieving definite outcomes that improve a patient’s quality of life. The outcomes include but are not limited to cure of a disease, elimination or reduction of a patient’s symptomatology, arresting or slowing of a disease process and the preventing of a disease or symptomatology.

(25) "Pharmacy internship" is a period of practical pharmacy experience under the direct supervision of a licensed pharmacist and pursuant to the rules of the board.

(26) "Pharmacy practice site" means any place within this state where prescription drugs or prescription devices are dispensed and where pharmaceutical care is provided, and any place outside of the state where prescription drugs or prescription devices are dispensed and pharmaceutical care is provided to persons residing in this state.

(27) "Preceptor" means an individual who is currently licensed as a pharmacist and who meets the qualifications of a preceptor under the rules of the board and participates in the education of pharmacy interns.

(28) "Prescription department" means the area of a pharmacy practice site in which prescription drugs and devices and related materials are stocked and medical and prescription orders are compounded and dispensed.

(29) "Quality assurance" means a system for identifying problems in patient care that are resolved via administrative, clinical, or educational actions to ensure that final products and outcomes meet applicable specifications.

(30) "Radiopharmaceutical service" means, but is not limited to:

(a) the compounding, dispensing, labeling, and delivering of radiopharmaceuticals;
(b) the participation in radiopharmaceutical selection and radiopharmaceutical utilization reviews;
(c) the proper and safe storage and distribution of radiopharmaceuticals;
(d) the maintenance of radiopharmaceutical quality assurance;
(e) the responsibility for advising, where necessary or where regulated, of the diagnostic and therapeutic value, hazards, and use of radiopharmaceuticals; and

(f) the offering or performing of those acts, services, operations, or transactions necessary in the conduct, operation, management, and control of a nuclear pharmacy practice site.

(31) "Reciprocity" means to issue a license to an applicant who furnishes satisfactory proof of licensing by examination in another state or territory pursuant to the rules of the board.

(32) "Shall" means that compliance is mandatory.

(33) "Sterile product" means any dosage form, drug product, or biological product devoid from all living microorganisms, including but not limited to bacteria and fungus.

(34) "Sterile manufacturing" means the production, propagation, processing, pooling, or repackaging of sterile products for wholesale or any other form of distribution, not pursuant to a prescription or medical order.

(35) "Third party pharmacy program" means any system of providing for the reimbursement of medical or prescription orders and/or pharmaceutical care services under a contractual arrangement or agreement between a provider of such services and the third party program administrator who is not the consumer of those services.

(36) "Third party pharmacy program administrator" means, but is not limited to, insurance companies, managed care organizations, health maintenance organizations, preferred provider organizations, pharmacy benefit managers, and pharmacy services administrative organizations.

(37) "Unit dose packaging" means that packaging which is designed to hold a quantity of a drug product intended for administration as a single dose.

(38) "USP" means the United States Pharmacopeia.

(39) "USP standards" means any applicable standard or standards published in the most current version of United States Pharmacopeia National Formulary guidelines, to the extent that such guidelines do not conflict with state law, rules, or Board Policy Statements and as those guidelines may, from time to time, be amended.


1140-01-02 VIOLATIONS CONSTITUTE UNPROFESSIONAL CONDUCT.

(1) Any person who violates any rule of the board may be deemed guilty of dishonorable, immoral, unethical or unprofessional conduct within the meaning of T.C.A. § 63-10-505(6).
Any person who violates any rule of the board may be deemed guilty of dishonorable, immoral, unethical or unprofessional conduct within the meaning of T.C.A. § 63-10-305(6).

Authority: T.C.A. §§ 63-10-101, 63-10-102, 63-10-202, 63-10-504(b)(1), and 63-10-505(6).


1140-01-.03 APPLICATION FOR A PHARMACIST LICENSE.

(1) An applicant for a license to engage in the practice of pharmacy shall submit the following to the Board office at time of application:

(a) A completed application on a form approved by the Board;

(b) Application and registration fees established in rule 1140-01-.10; and

(c) The result of a criminal background check, which the applicant shall pay for and cause to be submitted to the Board’s administrative office directly from the vendor identified in the Board’s licensure application materials.

(d) Any application submitted which lacks required information or reflects a failure to meet any of the requirements for licensure will be returned to the applicant with written notification of the information that is lacking or the reason(s) the application does not meet the requirements for licensure and will be held in “pending” status until satisfactorily completed within a reasonable period of time, not to exceed sixty (60) days from date of written notification.

(2) For the purpose of T.C.A. § 63-10-506(d), a “recognized” college or school of pharmacy is a college or school of pharmacy which meets the minimum standards of the ACPE and appears in the ACPE “Annual Directory of Accredited Professional Programs of Colleges and Schools of Pharmacy.”

(3) No applicant shall be eligible for a license if the applicant has engaged in conduct or suffers a condition which would constitute grounds for revocation or suspension of a license under T.C.A. § 63-10-505, unless the applicant can show cause why a license should be issued.

(4) No license shall be issued to a reciprocal applicant from a state which denies reciprocal privileges to a pharmacist currently licensed and in good standing in Tennessee.

(5) It shall be unlawful for any person to procure or attempt to procure a license or certificate of registration for such person or for any other person by making any false representations.

(6) An applicant initially licensed in another state and who wishes to obtain a Tennessee license may, in the discretion of the board, transfer to Tennessee the applicant's score on NAPLEX taken in another state. Provided, however, if the applicant has been licensed for twelve (12) or more months in another state, then the applicant shall apply for a license in Tennessee by reciprocity. No license shall be issued to a score transfer applicant from a state which denies score transfer privileges to a pharmacist currently licensed and in good standing in Tennessee.
1140-01-.04 PHARMACY INTERNSHIP.

(1) An applicant for an initial pharmacist license by examination must show, on affidavit forms prescribed by the board, that the applicant has acquired a minimum of one thousand five hundred (1,500) hours of pharmacy internship (practical pharmacy experience) under the instruction of a pharmacist in good standing, subject to all of the following conditions.

(a) The one thousand five hundred (1,500) hours must be acquired after enrollment in a recognized college or school of pharmacy; one thousand one hundred (1,100) of these hours may be acquired in pharmacy programs or demonstration projects structured by the college or school of pharmacy.

(b) Pharmacy internship may be acquired in another state, provided that the preceptor's qualifications are certified by the appropriate authorities of such state.

(c) Four hundred (400) of these hours may be acquired in non-traditional pharmacy internship programs which have received prior approval of the board.

(d) Foreign pharmacy graduates shall complete five hundred (500) hours of pharmacy internship in Tennessee within a period of six (6) consecutive months.


1140-01-.05 LICENSING EXAMINATIONS.

(1) An applicant for an initial license to engage in the practice of pharmacy in the State of Tennessee shall take the National Association of Boards of Pharmacy (NABP) Multistate Pharmacy Jurisprudence Examination (MPJE®) and the NABP North American Pharmacy Licensing Examination (NAPLEX®), which shall be administered on the dates scheduled by the NABP. An applicant shall also meet the minimum acceptable passing scores on the NAPLEX® and MPJE® as established and nationally accepted.

(2) An applicant to obtain a pharmacy license by reciprocity shall successfully complete the MPJE®.

(3) In addition to completing the requirements in paragraph (1) of this rule, a pharmacy foreign graduate shall successfully complete the foreign pharmacy equivalency examination, the Test of Spoken English (TSE®) examination and any other requirements established by the NABP.

(4) Any applicant who fails either the NAPLEX® or MPJE® may retake the examinations at any one of the next examination dates scheduled by the NABP. If an applicant fails the NAPLEX® or MPJE® three (3) consecutive times, then the Board may require that applicant to take review courses prior to any following reexamination.

1140-01-.06 SUMMARY SUSPENSION OF LICENSE.

Pursuant to T.C.A. § 4-5-320, if the board finds that public health, safety or welfare imperatively requires emergency action and incorporates a finding to that effect in its order, summary suspension of a license may be ordered pending proceedings for revocation or other action.

Authority: T.C.A. §§ 4-5-320, 63-10-101, 63-10-102, 63-10-504(b)(1), 63-10-504(b)(2), and 63-10-505.


1140-01-.07 INACTIVE LICENSES AND LICENSE REINSTATEMENT.

(1) A pharmacist may apply for an inactive license by:

(a) Completing the biennial license renewal application form; and

(b) Paying the biennial renewal fee for an inactive license.

(2) A pharmacist maintaining an active license to practice pharmacy in another state or jurisdiction is ineligible for inactive license status in Tennessee.

(3) A pharmacist seeking active status for an inactive, delinquent, suspended or revoked license must fulfill the following minimum requirements.

(a) If the license has been inactive, delinquent, suspended or revoked for less than one (1) year, the pharmacist shall:

1. Provide written notice to the board requesting an active license;

2. Satisfy all past due continuing pharmaceutical education as required by the board;

3. Pay all cumulative license renewal fees and any applicable penalty fees for the period during which the license was inactive, delinquent, suspended or revoked.

(b) If the license has been inactive, delinquent, suspended or revoked from one (1) year to not more than five (5) consecutive years, the pharmacist shall:

1. Provide written notice to the board requesting an active license;

2. Satisfy all past due continuing pharmaceutical education as required by the board;

3. Successfully complete the jurisprudence examination;

4. Pay all cumulative license renewal fees and any applicable penalty fees for the period during which the license was inactive, delinquent, suspended or revoked; and

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5. Complete a period of pharmacy internship in Tennessee as follows.

   (i) If the license has been inactive, delinquent, suspended or revoked from one (1) year to not more than three (3) consecutive years, one hundred sixty (160) hours within ninety (90) consecutive days.

   (ii) If the license has been inactive, delinquent, suspended or revoked for more than three (3) consecutive years but not more than five (5) consecutive years, three hundred twenty (320) hours within one hundred eighty (180) consecutive days.

(c) If the license has been inactive, delinquent, suspended or revoked for more than five (5) consecutive years, the pharmacist shall:

1. Provide written notice to the board requesting an active license;
2. Satisfy all past due continuing pharmaceutical education as required by the board;
3. Successfully complete the NAPLEX and jurisprudence examinations;
4. Pay all cumulative license renewal fees and any applicable penalty fees for the period during which the license was inactive, delinquent, suspended or revoked; and
5. Complete a period of pharmacy internship of three hundred twenty (320) hours within one hundred eighty (180) consecutive days.

(d) Fulfill any other requirements which may be contained in any order of the board suspending or revoking the applicant's license.

(e) The board shall consider a written notice requesting reinstatement of an inactive, delinquent, suspended or revoked license within ninety (90) days of the notice being received by the director.

(f) The board shall consider a waiver upon request.

Authority:  T.C.A. §§ 63-10-101, 63-10-102, 63-10-210, 63-10-404(17), and 63-10-504(b)(1).

1140-01-.08 APPLICATION FOR PHARMACY PRACTICE SITE, MANUFACTURER AND WHOLESALER/DISTRIBUTOR LICENSES.

(1) Application for a license to operate as a pharmacy practice site, manufacturer or wholesaler/distributor within the state of Tennessee shall be submitted to the office of the board at least thirty (30) days prior to the scheduled opening date. No pharmacy practice site, manufacturer or wholesaler/distributor may open within the state of Tennessee until a license has been obtained; and such license will not be issued until an inspection by an authorized representative of the board has been made.

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(2) An application for an existing pharmacy practice site, manufacturer or wholesaler/distributor physically located within the state of Tennessee must be filed when the pharmacy practice site, manufacturer or wholesaler/distributor changes name, location or ownership.

(a) Transactions constituting a change of ownership include, but are not limited to, the following:

1. A sole proprietor becomes a member of a partnership or corporation, which succeeds him as the new operator;
2. A partnership dissolves;
3. One partnership is replaced by another through the removal, addition or substitution of a partner;
4. Two (2) or more corporations merge and the originally-licensed corporation does not survive; and
5. Transfers between levels of government.

(b) Transactions which do not constitute a change of ownership include, but are not limited to, the following:

1. Changes in the membership of a corporate board of directors or board of trustees;
2. Two (2) or more corporations merge and the originally-licensed corporation survives; and
3. Corporate stock transfers or sales, even when a controlling interest.

(3) No out-of-state pharmacy practice site, manufacturer or wholesaler/distributor shall conduct business in the state of Tennessee until such pharmacy practice site, manufacturer or wholesaler/distributor obtains the required license from the board. In order to obtain a license for a pharmacy practice site, manufacturer or wholesaler/distributor physically located out-of-state the following standards must be met.

(a) Pharmacy practice site.

1. Submit an application for a license, which shall include the address of the pharmacy practice site, name of owner if a sole proprietorship, names of partners if a partnership or names and titles of all officers if a corporation and names of all pharmacists who practice at the site, together with the appropriate application fee. The director shall be notified in writing within thirty (30) days of any change in the information contained on the original application for a license, including names of pharmacists practicing at the site.
2. Comply with all statutorily authorized directions and requests for information from the board.
3. Maintain at all times a current permit, license or registration to conduct the pharmacy practice site in compliance with the laws of the state in which the site is physically located.
4. Submit a copy of the most recent inspection report resulting from an inspection conducted by the regulatory or licensing agency of the state in which the pharmacy practice site is physically located. Thereafter, the pharmacy practice site shall submit to the director a copy of any subsequent inspection report conducted by the regulatory or licensing agency of the state in which the site is physically located.

5. Maintain records of prescription orders dispensed to persons residing in Tennessee.

6. All records of prescription orders prepared and dispensed to persons residing in Tennessee shall be readily retrievable from other records.

7. During regular hours of operation, but not less than six (6) days per week nor for a minimum of forty (40) hours per week provide access to a pharmacist by a toll-free telephone service. A toll-free number shall be placed on the label affixed to the dispensing container for each prescription dispensed to a person residing in Tennessee.

8. Designate a pharmacist in charge who shall be responsible for compliance with the provisions in this section, and who shall hold a current Tennessee pharmacist license.

9. All out-of-state pharmacy practice sites shall comply with the requirements for patient counseling, patient profiling, drug regimen review and pharmaceutical care as set forth at 1140-03-.01.

(b) Manufacturer or wholesaler/distributor.

1. Submit an application for a license, which shall include the address of the manufacturer or wholesaler/distributor, name of owner if a sole proprietorship, names of partners if a partnership or names and titles of all officers if a corporation, together with the appropriate application fee. The director shall be notified in writing within thirty (30) days of any change in the information contained on the original application for a license.

2. Submit a copy of the most recent inspection report resulting from an inspection conducted by the regulatory or licensing agency of the state in which the manufacturer or wholesaler/distributor is physically located. Thereafter, the manufacturer or wholesaler/distributor shall submit to the director a copy of any subsequent inspection report conducted by the regulatory or licensing agency of the state in which the manufacturer or wholesaler/distributor is physically located.

3. Comply with the requirements contained in Chapter 1140-09 of the rules of the Board of Pharmacy.

(4) Representatives of a manufacturer or wholesaler/distributor conducting business in the state of Tennessee and who possess and distribute controlled substances shall obtain a controlled substance registration from the Board of Pharmacy.

(5) A manufacturer conducting business in the state of Tennessee and who manufactures, prepares, propagates, repackages, or processes sterile drug products or biological products using aseptic processing must register and possess a modifier as a sterile manufacturer with the Board of Pharmacy in accordance with this chapter. This section shall not apply to wholesalers/distributors of sterile products.
It shall be unlawful for any person to procure or attempt to procure a license or certificate of registration for such person or for any other person by making any false representations.

In determining whether to grant a license under this rule, the board shall require from the applicant proof satisfactory to the board that the:

(a) Applicant is of good moral character, or, if the applicant is a partnership or corporation, that the managing officers are of good moral character; and

(b) That the applicant is equipped as to land, buildings and equipment necessary to conduct the business for which the application has been submitted.


1140-01-.09 RENEWAL OF LICENSES.

(1) All licenses and certificates of registration granted by the board shall be for a two (2) year period beginning on the date the license is initially granted. All licenses and certificates of registration shall be renewed on or before the last day of the two (2) year license cycle.

(2) A pharmacist or pharmacy technician serving in the uniformed services of the United States shall not be required to pay license or registration renewal fees during the period of active duty and the pharmacist shall not be required to complete continuing pharmacy education requirements during the period of active duty.


1140-01-.10 FEES.

(1) An applicant for examination for a license as a pharmacist shall pay a fee of fifty dollars ($50.00) plus cost of the examination and materials.

(2) An applicant for a reciprocal license or NAPLEX score transfer shall pay a fee of three hundred dollars ($300.00).

(3) Each person becoming licensed as a pharmacist shall pay a registration fee of one-hundred twenty-five dollars ($125.00). Each person licensed as a pharmacist who desires to continue in the practice of pharmacy shall biennially, on or before the last day of the month that the person’s license shall expire, pay a renewal fee of one-hundred twenty-five dollars ($125.00). Each person licensed as a pharmacist and who wishes to obtain an inactive license shall biennially, on or before the last day of the month that the person’s license shall expire, pay a renewal fee of sixty-three dollars ($63.00).
(4) Each person becoming registered as a pharmacy technician shall pay a registration fee of 
seventy-five dollars ($75.00). Each person who desires to continue to practice as a 
pharmacy technician shall biennially, on or before the last day of the month that the person’s 
registration shall expire, pay a renewal fee of seventy-five dollars ($75.00).

(5) Any person, partnership, firm, corporation or agency owning or operating a pharmacy 
practice site or any establishment or institution where prescription drugs and devices and 
related materials are kept for the purpose of the compounding and dispensing of medical and 
prescription orders shall pay a registration fee of three-hundred dollars ($300.00) biennially. 
Any new pharmacy practice site to be opened or established, or any change in location, 
name or ownership of any existing pharmacy practice site, shall before active operation 
obtain a license from the Board of Pharmacy and shall pay a fee of three-hundred dollars 
($300.00).

(6) All manufacturers and wholesalers/distributors of prescription drugs and devices and related 
materials doing business in the state of Tennessee must be licensed by the Board of 
Pharmacy by paying a registration fee of five-hundred twenty-five dollars ($525.00), and 
thereafter a biennial renewal fee of five-hundred twenty-five dollars ($525.00).

(7) The fee for the Board of Pharmacy’s publication of Pharmacy Drug Laws, Rules and 
Regulations shall be an amount which covers the cost of publication and shipping, as 
determined by the Board of Pharmacy.

(8) The charge for a roster of Tennessee pharmacies, pharmacists and printing of mailing labels 
of Tennessee pharmacies and pharmacists shall be determined by the administration of the 
Department of Health.

(9) The fee for certification of license examination grades shall be twenty five dollars ($25.00).

(10) The fee for a duplicate or revised pharmacist license wall certificate shall be twenty five 
dollars ($25.00).

(11) If any person fails to renew a license, such license may be reinstated upon complying with 
rule 1140-01-.07 and upon the payment of the appropriate renewal fee plus a penalty fee of 
ten dollars ($10.00) for each month or fraction thereof that payment for renewal is delinquent. 
In the event such renewal is not procured within six (6) months from the date on which the 
last renewal became delinquent, the board may refuse to issue the renewal.

(12) If any person fails to renew a license or registration certificate, such license or registration 
certificate may be reinstated upon complying with rule 1140-01-.07 and upon the payment of 
the appropriate renewal fee plus a penalty fee of ten dollars ($10.00) for each month or 
fraction thereof that payment for renewal is delinquent. In the event such renewal is not 
procured within six (6) months from the date on which the last renewal became delinquent, 
the board may refuse to issue the renewal.

(13) A penalty of fifty dollars ($50.00) may, in the discretion of the board, attach to each failure of 
a licensee or registration certificate holder to provide any required notice to the director as 
may be required by the rules of the board.

(14) Any licensee who wishes to modify the terms or conditions of a license to manufacture, 
obtain, possess, administer or dispense a prescription drug or device or controlled substance 
for the purpose of scientific research, chemical analysis, instruction or training of detection 
animals shall file those modifications with a non-refundable fee of five dollars ($5.00).
Any person who holds a license to manufacture, obtain, possess, administer or dispense a prescription drug or device or controlled substance for the purpose of scientific research, chemical analysis, instruction or training of detection animals shall pay a renewal fee of one-hundred dollars ($100.00) biennially from the date of issuance.

Any person, partnership, firm, corporation or agency owning or operating a pharmacy practice site or any other establishment licensed pursuant to this chapter, where sterile products are compounded, manufactured, prepared, propagated, repackaged, processed, stored, or distributed shall pay a registration fee of two-hundred and fifty dollars ($250.00), and thereafter a biennial renewal fee of two-hundred and fifty dollars ($250.00).


CONTROLLED SUBSTANCE REGISTRATION.

No licensee may obtain, possess, administer, dispense, distribute, or manufacture any controlled substance in this state, and no representative of a manufacturer or wholesaler/distributor may distribute any controlled substance in this state, without obtaining a controlled substance registration from the board. Application for such registration shall be submitted on a form prescribed by the board, and shall be accompanied by a fee of forty dollars ($40.00) and thereafter a biennial renewal fee of forty dollars ($40.00).


STERILE COMPOUNDING REGISTRATION

(1) No licensee may compound, manufacture, prepare, propagate, or process any sterile product to be dispensed, sold, traded, or otherwise distributed in or from this state without first obtaining a sterile compounding modifier registration from the Board of Pharmacy.

(2) A registration modifier to compound and dispense sterile products into or from this state may be suspended by the Board of Pharmacy, upon information that the registrant has:

(a) Knowingly furnished false or fraudulent material information in any application filed before the Board of Pharmacy; or

(b) Been convicted of a felony under any state or federal law relating to drugs or to the practice of pharmacy; or

(c) Had any of its licenses, permits, or registrations granted by the Food and Drug Administration (FDA), the Drug Enforcement Administration (DEA), the Department of Health and Human Services (DHHS), or any other federal agency or subdivision thereof, suspended, revoked, or voluntarily surrendered; or

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(d) Been enjoined from operation by the court of any state or a federal court; or

(e) Been identified by the Commissioner of Health or the Commissioner’s designee, the Centers for Disease Control and Prevention (CDC), the Food and Drug Administration (FDA), or an investigator of the Board of Pharmacy as a source of adulterated, misbranded, or otherwise unsafe sterile products which have been, or pose an imminent risk of being dispensed, sold, traded, or otherwise distributed.

(3) An order of suspension issued by the Board of Pharmacy may contain additional directives or requirements necessary to protect public health, safety and welfare, including but not limited to:

(a) The quarantine or disposal of any sterile product compounded, manufactured, prepared, propagated or processed at the facility.

(b) The initiation of a recall of any sterile product compounded, manufactured, prepared, propagated or processed at the facility where such products or any label, container, packaging, or dosage form associated with such products may be adulterated, misbranded, contaminated, or otherwise unsafe.

(c) An order of suspension issued by the Board of Pharmacy may contain exceptions or allowances necessary to protect individual patients or the public health.

(4) Any order of suspension issued by the Board of Pharmacy pursuant to this chapter shall follow the procedures required by the Uniform Administrative Procedures Act, including those procedures required by T.C.A. § 4-5-320(d) where appropriate.

1140-01-.13 1140-01-.13, STANDARDS FOR PHARMACIES AND PRESCRIPTION DEPARTMENT SECURITY.

A license to operate a new or remodeled pharmacy practice site, or an existing pharmacy practice site which changes location or ownership, will not be issued unless the pharmacy practice site meets the following standards.

(1) The pharmacy practice site and equipment therein shall be maintained in a clean, sanitary, orderly and well-lighted condition, and all persons working in the pharmacy practice site shall be required to keep themselves and their apparel in a clean and sanitary condition.

(2) All new or relocated pharmacies opening after July 1, 1998 shall provide a consultation area which offers sufficient privacy to the patient before a license will be issued. All existing pharmacies shall be in compliance with this requirement on or before January 1, 2000.

(3) The prescription department at the pharmacy practice site shall meet the following standards.

(a) The department shall have necessary counters and storage space.

(b) The department shall have a representative stock of prescription drugs and devices and related materials sufficient to compound and dispense medical and prescription orders as indicated by experience.

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(c) The department shall have the apparatus and equipment needed to compound and dispense medical and prescription orders properly.

(d) The department shall occupy a space of not less than one hundred eighty (180) square feet.

(e) The department shall have hot and cold running water and immediate area refrigeration.

(f) The department shall have a physical barrier sufficient to protect against unauthorized entry and pilferage of prescription drugs and devices and related materials.

(g) Keys or other access devices to the physical barriers shall be subject to the following standards.

1. Only pharmacists practicing at the pharmacy and pharmacists authorized by the pharmacist in charge shall be in possession of any keys or other access devices.

2. The pharmacist in charge shall place a key or other access device in a sealed envelope bearing the signature of the pharmacist in charge affixed across the seal and placed in a safe or vault in a secured place outside of the department. The key or access device may be used to allow emergency entrance to the department.

(h) Access to the department is restricted to pharmacists, pharmacy interns and pharmacy technicians who are practicing at the pharmacy. Other persons designated by the pharmacist in charge may be allowed access but only during hours that a pharmacist is on duty.

(i) Notwithstanding any rule or regulation to the contrary, a pharmacy which was established before June 6, 1945, and which serves food, and which has continuously had a soda fountain, may allow a customer to go through the pharmacy area to the restroom, and not be required to have a gate or door to separate the pharmacy from the restroom or other parts of the establishment.

(4) All licenses and certificates of registration for a pharmacy practice site shall at all times be conspicuously displayed at the practice site.

(5) If a pharmacy practice site is located in a mercantile establishment (such as a discount store, grocery store, department store, or other similar establishment), then such pharmacy practice site shall be:

(a) open for business during the same hours as the mercantile establishment, unless the pharmacy practice site is capable of being closed-off by physical barrier from floor to ceiling; and

(b) under the supervision of a pharmacist at all times; except as provided in rule 1140-03-.07.

(6) The pharmacist shall not at any time be denied access to the prescription department of a pharmacy practice site located in a mercantile establishment; provided, however, that entry of the pharmacist at times when the pharmacy is closed to the public may be subject to reasonable and prudent conditions.

(7) A pharmacy practice site where prescription drugs and devices and related materials are received, stored, compounded and dispensed shall not be opened for business or any other
(Rule 1140-01-.12, continued)

reason unless a licensed pharmacist is present. Furthermore, no medical or prescription
order shall be dispensed except during the presence and under the direct supervision of a
pharmacist.

(8) Nothing in this rule applies to a pharmacy practice site or prescription department operating
in an institutional facility.

(9) In cases of practical difficulty or undue hardship, the board may permit exceptions to the
standards specified in this rule.

Authority: T.C.A. §§ 63-10-404(28), 63-10-504(b)(1), and 63-10-504(b)(1) and (2). Administrative

1140-01-.14

STANDARDS FOR MANUFACTURERS AND
WHOLESALER/DISTRIBUTORS.

No license to operate a new or remodeled manufacturer or wholesaler/distributor location within the state
of Tennessee, or an existing manufacturer or wholesaler/distributor location which changes location or
ownership, will be issued unless the manufacturer or wholesaler/distributor meets the standards set forth
in Chapter 1140-09 of the rules of the Board of Pharmacy.

Authority: T.C.A. §§ 63-10-404(18) and (37), and 63-10-504(b)(1). Administrative History: Original

1140-01-.15

PRESCRIPTION DRUGS DISPENSED BY HEALTH DEPARTMENTS.

For purposes of T.C.A. § 63-10-405, the following drugs are hereby approved as not subject to abuse:

(1) Tuberculosis Control Agents:

(a) Capreomycin Injection
(b) Cycloserine Capsules
(c) Ethambutol Tablets
(d) Ethionamide Tablets
(e) Isoniazid Tablets
(f) Para-Aminosalicylate Tablets
(g) Pyrazinamide Tablets
(h) Rifampin Capsules
(i) Streptomycin Injection
(j) Tuberculin Skin Test (Mantoux only)
(k) Rifampin/Isoniazid
(l) Ofloxacin
(m) Rifampin-Isoniazid-Pyrazinamide

(2) Venereal Disease Control Agents:

(a) Ampicillin Capsules
(b) Doxycycline Capsules
(c) Erythromycin Tablets
(d) Penicillin

1. Benzathine Penicillin G Injection
2. Procaine Penicillin G Injection

(e) Probenecid Tablets
(f) Spectinomycin Injection
(Rule 1140-01-.14, continued)

(g) Tetracycline Capsules
(h) Ceftriaxone
(i) Ciprofloxacin
(j) Lidocaine Injection
(k) Azithromycin
(l) Acyclovir Tablets, Ointments
(m) Trichloroacetic Acid
(n) Salicylic Acid
(o) Podophyllin/Salicylic Acid
(p) Aldara (Imiquimod)

(3) Biologicals/Immunizations:

(a) Antisera
(b) Antitoxins
(c) Immune Serum Globulin
(d) Toxoids
(e) Vaccines
(f) Antigens

(4) Reproductive Health Agents:

(a) Metronidazole Tablets
(b) Oral Contraceptives
(c) Podophyllin
(d) Prenatal Vitamins
(e) Triple Sulfa Vaginal Cream/Tabs
(f) Vaginal Antifungal Cream/Tabs

1. Clotrimazole
2. Miconazole
3. Nystatin
4. Terconazole (Terazole)

(g) Amino-Cerv
(h) Nitrofurantoin
(i) Ibuprofen, 600 mg Tablets
(j) Metronidazole (vaginal jell)
(k) Fluconazole Tablets
(l) Clindamycin Vaginal Cream
(m) Premarin Tablets (for use in estrogen trials for the evaluation of atypical cells in certain inflammatory atrophic pap smears)
(n) Medroxyprogesterone Acetate Injectable (Depo Provera®)
(o) Norelgestromin/ethinyl estradiol transdermal system (Ortho Evra®)

(p) Etonogestrel/ethinyl estradiol vaginal ring (Nuvaring®)

(5) Child Health Agents:

(a) Fluoride Tablets and Drops
(b) Lindane Cream, Lotion, Shampoo
(c) Mebendazole Tablets
(d) Pyrantel Pamoate Liquid
(e) Sulfadiazone Tablets
(f) Trimethoprim and Sulfamethoxazole
(g) Permethrin

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(Rule 1140-01-.14, continued)

(h) Crotamiton

(i) Nystatin Oral Suspension

(j) Nystatin Triamcinolone Cream

(k) Ibuprofen, Suspension Liquid

(6) Emergency Agents:

(a) Aminophylline Injection

(b) Benztropine Injection

(c) Diphenhydramine Injection

(d) Epinephrine Injection

(e) Glucagon Injection

(f) Hydralazine Injection

(g) Hydrocortisone Sodium Succinate

(h) Insulin, Regular

(i) Intravenous Fluids

(j) Oxygen

(k) Phenylephrine Injection

(l) Sodium Bicarbonate Injection

(m) Atropine Injection

(n) Nitroglycerin Sublingual Tablets

(o) Dexamethasone Injection

(p) Norepinephrine

(7) Antihypertensive Agents:

(a) Methyldopa

(b) Reserpine

(c) Hydrochlorothiazide

(d) Hydralazine

(e) Propranolol

(f) Potassium Supplements

(g) Nicotine Patches