Virginia Board of Pharmacy Prescriptive Authority in Virginia

Reference: § 54.1-3400 *et seq*. of the Code of Virginia commonly known as the Drug Control Act and § 54.1-3303 of the Code of Virginia, and respective Board regulations.

In Virginia all prescription drugs are categorized into schedules. Schedules I through V, for the most part, mirror the federal schedules. All <u>prescription</u> or <u>legend</u> drugs not included in Schedules II through V are placed in Schedule VI in Virginia and are also referred to as "controlled" drugs or substances within the Drug Control Act. This is sometimes confusing as the term "controlled" is usually applied only to drugs in Schedules II through V.

Before prescribing any drug in Schedules II-V, a practitioner must obtain a registration from the U.S Drug Enforcement Administration (DEA). The DEA registration must also be on any prescription written for a Schedule II-V drug.

Nurse practitioners who meet certain criteria may be issued a license to prescribe Schedule II-VI drugs by the Boards of Nursing and Medicine. Authorization to prescribe schedules or categories of drugs will be set out in a practice agreement with a collaborating physician. Nurse practitioners with prescriptive authority may dispense samples of those drugs they are authorized to prescribe and may also sign for the receipt of those samples.

Physician assistants (PA's) who meet criteria and have been approved by the Board of Medicine for prescriptive authority may prescribe Schedule II-VI drugs that have been approved by the supervising medical practitioner or podiatrist. A prescription written by a physician assistant for a Schedule II-V drug must include the name of the supervising physician. Physician assistants may dispense samples of those drugs they are authorized to prescribe and may sign for receipt of samples.

Nurse practitioners or physician assistants whose prescriptive authority is limited to Schedule VI are not legally required to have a DEA number but will possess a Virginia license. For nurse practitioners, there is a 10-digit license number beginning with 0017, which should be on the prescription and can be verified through the web site <u>www.dhp.virginia.gov</u> under "on-line license lookup" and checking the occupation "Authorization to Prescribe." For physician assistants, there is a 10-digit license number beginning with 011, which can be verified through the web site <u>www.dhp.virginia.gov</u> under "on-line license lookup" and checking the occupation "Authorization to Prescribe." For physician assistants, there is a 10-digit license number beginning with 011, which can be verified through the web site <u>www.dhp.virginia.gov</u> under "on-line license lookup" and checking the occupation "Physician Assistant."

Practitioners of medicine, osteopathy, podiatry, dentistry, or veterinary medicine have independent prescriptive authority and may prescribe drugs in Schedules II through VI.

Optometrists who have been certified to use therapeutic pharmaceutical agents have independent authority to prescribe and administer certain controlled substances and devices to treat diseases and abnormal conditions of the human eye and its adnexa in these categories:

1. Oral analgesics - Schedule II controlled substances consisting of hydrocodone in combination with acetaminophen and Schedule III, IV and VI narcotic and non-narcotic agents.

- 2. Topically administered Schedule VI agents:
 - a. Alpha-adrenergic blocking agents;
 - b. Anesthetic (including esters and amides);
 - c. Anti-allergy (including antihistamines and mast cell stabilizers);
 - d. Anti-fungal;

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- e. Anti-glaucoma (including carbonic anhydrase inhibitors and hyperosmotics);
- f. Anti-infective (including antibiotics and antivirals);
- g. Anti-inflammatory;
- h. Cycloplegics and mydriatics;
- i. Decongestants; and
- j. Immunosuppressive agents.
- 3. Orally administered Schedule VI agents:
 - a. Aminocaproic acids (including antifibrinolytic agents);
 - b. Anti-allergy (including antihistamines and leukotriene inhibitors);
 - c. Anti-fungal;
 - d. Anti-glaucoma (including carbonic anhydrase inhibitors and hyperosmotics);
 - e. Anti-infective (including antibiotics and antivirals);
 - f. Anti-inflammatory (including steroidal and non-steroidal);
 - g. Decongestants; and
 - h. Immunosuppressive agents.

Inquiries as to the certification of an optometrist to prescribe therapeutic pharmaceutical agents or requests for regulations may be made by checking the web site <u>www.dhp.virginia.gov</u> under "on-line license lookup" and checking for the occupation "TPA certified optometrist." After June 30, 2004, every person who is initially licensed to practice optometry in Virginia must meet the qualifications for a TPA-certified optometrist.

In order to be valid, prescriptions must meet the criteria set forth in § 54.1-3303 of the Code of Virginia (attached). A prescription must be written in the context of a bona fide practitioner-patient relationship, for a medicinal or therapeutic purpose, and within the course of the professional practice of the prescriber. The elements that constitute a bona fide practitioner patient relationship are set forth in this statute.

from the Code of Virginia:

A. A prescription for a controlled substance may be issued only by a practitioner of medicine, osteopathy, podiatry, dentistry or veterinary medicine who is authorized to prescribe controlled substances, or by a licensed nurse practitioner pursuant to 54.1-2957.01, a licensed physician assistant pursuant to 54.1-2952.1, or a TPA-certified optometrist pursuant to Article 5 (54.1-3222 et seq.) of Chapter 32. The prescription shall be issued for a medicinal or therapeutic purpose and may be issued only to persons or animals with whom the practitioner has a bona fide practitioner-patient relationship.

For purposes of this section, a bona fide practitioner-patient-pharmacist relationship is one in which a practitioner prescribes, and a pharmacist dispenses, controlled substances in good faith to his patient for a medicinal or therapeutic purpose within the course of his professional practice. In addition, a bona fide practitioner-patient relationship means that the practitioner shall (i) ensure that a medical or drug history is obtained; (ii) provide information to the patient about the benefits and risks of the drug being prescribed; (iii) perform or have performed an appropriate examination of the patient, either physically or by the use of instrumentation and diagnostic equipment through which images and medical records may be transmitted electronically; except for medical emergencies, the examination of the patient shall have been performed by the practitioner himself, within the group in which he practices, or by a consulting practitioner prior to issuing a prescription; and (iv) initiate additional interventions and follow-up care, if necessary, especially if a prescribed drug may have serious side effects.

For the purpose of prescribing a Schedule VI controlled substance to a patient via telemedicine services as defined in § <u>38.2-3418.16</u>, a prescriber may establish a bona fide practitioner-patient relationship by an examination through face-to-face interactive, two-way, real-time communications services or store-and-forward technologies when all of the following conditions are met: (a) the patient has provided a medical history that is available for review by the prescriber; (b) the prescriber obtains an updated medical history at the time of prescribing; (c) the prescriber makes a diagnosis at the time of prescribing; (d) the prescriber conforms to the standard of care expected of in-person care as appropriate to the patient's age and presenting condition, including when the standard of care requires the use of diagnostic testing and performance of a physical examination, which may be carried out through the use of peripheral devices

Revised: January 15, 2016

appropriate to the patient's condition; (e) the prescriber is actively licensed in the Commonwealth and authorized to prescribe; (f) if the patient is a member or enrollee of a health plan or carrier, the prescriber has been credentialed by the health plan or carrier as a participating provider and the diagnosing and prescribing meets the qualifications for reimbursement by the health plan or carrier pursuant to § <u>38.2-3418.16</u>; and (g) upon request, the prescriber provides patient records in a timely manner in accordance with the provisions of § <u>32.1-127.1:03</u> and all other state and federal laws and regulations. Nothing in this paragraph shall permit a prescriber to establish a bona fide practitioner-patient relationship for the purpose of prescribing a Schedule VI controlled substance when the standard of care dictates that an in-person physical examination is necessary for diagnosis. Nothing in this paragraph shall apply to: (1) a prescriber providing on-call coverage per an agreement with another prescriber or his prescriber's professional entity or employer; (2) a prescriber consulting with another prescriber regarding a patient's care; or (3) orders of prescribers for hospital out-patients or in-patients.

Any practitioner who prescribes any controlled substance with the knowledge that the controlled substance will be used otherwise than medicinally or for therapeutic purposes shall be subject to the criminal penalties provided in § <u>18.2-248</u> for violations of the provisions of law relating to the distribution or possession of controlled substances.

B. In order to determine whether a prescription that appears questionable to the pharmacist results from a bona fide practitioner-patient relationship, the pharmacist shall contact the prescribing practitioner or his agent and verify the identity of the patient and name and quantity of the drug prescribed. The person knowingly filling an invalid prescription shall be subject to the criminal penalties provided in § <u>18.2-248</u> for violations of the provisions of law relating to the sale, distribution or possession of controlled substances. No prescription shall be filled unless there is a bona fide practitioner-patient-pharmacist relationship. A prescription not issued in the usual course of treatment or for authorized research is not a valid prescription.

C. Notwithstanding any provision of law to the contrary and consistent with recommendations of the Centers for Disease Control and Prevention or the Department of Health, a practitioner may prescribe Schedule VI antibiotics and antiviral agents to other persons in close contact with a diagnosed patient when (i) the practitioner meets all requirements of a bona fide practitioner-patient relationship, as defined in subsection A, with the diagnosed patient; (ii) in the practitioner's professional judgment, the practitioner deems there is urgency to begin treatment to prevent the transmission of a communicable disease; (iii) the practitioner has met all requirements of a bona fide practitioner-patient relationship, as defined in subsection A, for the close contact except for the physical examination required in clause (iii) of subsection A; and (iv) when such emergency treatment is necessary to prevent imminent risk of death, life-threatening illness, or serious disability.

D. A pharmacist may dispense a controlled substance pursuant to a prescription of an out-of-state practitioner of medicine, osteopathy, podiatry, dentistry or veterinary medicine authorized to issue such prescription if the prescription complies with the requirements of this chapter and the Drug Control Act (§ <u>54.1-3400</u> et seq.).

E. A licensed nurse practitioner who is authorized to prescribe controlled substances pursuant to $\frac{54.1-2957.01}{54.1-2957.01}$ may issue prescriptions or provide manufacturers' professional samples for controlled substances and devices as set forth in the Drug Control Act ($\frac{54.1-3400}{54.1-3400}$ et seq.) in good faith to his patient for a medicinal or therapeutic purpose within the scope of his professional practice.

F. A licensed physician assistant who is authorized to prescribe controlled substances pursuant to $\frac{54.1-2952.1}{1000}$ may issue prescriptions or provide manufacturers' professional samples for controlled substances and devices as set forth in the Drug Control Act ($\frac{54.1-3400}{1000}$ et seq.) in good faith to his patient for a medicinal or therapeutic purpose within the scope of his professional practice.

G. A TPA-certified optometrist who is authorized to prescribe controlled substances pursuant to Article 5 (§ 54.1-3222 et seq.) of Chapter 32 may issue prescriptions in good faith or provide manufacturers' professional samples to his patients for medicinal or therapeutic purposes within the scope of his professional practice for the drugs specified on the TPA-Formulary, established pursuant to § 54.1-3223, which shall be limited to (i) analgesics included on Schedule II controlled substances as defined in § 54.1-3448 of the Drug Control Act (§ 54.1-3400 et seq.) consisting of hydrocodone in combination with acetaminophen; (ii) oral analgesics included in Schedules III through VI, as defined in §§ 54.1-3450 and 54.1-3455 of the Drug Control Act (§ 54.1-3400 et seq.), which are appropriate to relieve ocular pain; (iii) other oral Schedule VI controlled substances, as defined in § 54.1-3455 of the Drug Control Act, appropriate to treat diseases and abnormal conditions of the human eye and its adnexa; (iv) topically applied Schedule VI drugs, as defined in § 54.1-3455 of the Drug Control Act; and (v) intramuscular administration of epinephrine for treatment of emergency cases of anaphylactic shock.

Revised: January 15, 2016

H. The requirement for a bona fide practitioner-patient relationship shall be deemed to be satisfied by a member or committee of a hospital's medical staff when approving a standing order or protocol for the administration of influenza vaccinations and pneumococcal vaccinations in a hospital in compliance with 32.1-126.4.