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# Notice of Intended Regulatory Action (NOIRA) Agency Background Document

| Agency name   | Board of Pharmacy, Department of Health Professions            |
|---|--|
| Virginia Administrative Code<br>(VAC) Chapter citation(s) | 18VAC110-20  |
| VAC Chapter title(s)                                      | Regulations Governing the Practice of Pharmacy                 |
| Action title  | Allowance for centralized pharmacy to verify Schedule VI drugs |
| Date this document prepared                               | 10/7/21  |

This information is required for executive branch review and the Virginia Registrar of Regulations, pursuant to the Virginia Administrative Process Act (APA), Executive Order 14 (as amended, July 16, 2018), the Regulations for Filing and Publishing Agency Regulations (1VAC7-10), and the *Form and Style Requirements for the Virginia Register of Regulations and Virginia Administrative Code*.

### **Brief Summary**

Provide a brief summary (preferably no more than 2 or 3 paragraphs) of the subject matter, intent, and goals of this this regulatory change (i.e., new regulation, amendments to an existing regulation, or repeal of an existing regulation.

In response to a petition for rulemaking, the Board is issuing a Notice of Intended Regulatory Action to amend sections 460 and 490 to allow a pharmacist at a central distribution company to verify Schedule VI drugs to be placed in an ADD prior to delivery to the receiving hospital and pharmacy technicians at the hospital to load the drugs directly into the ADD without further verification by a pharmacist at the hospital.

### **Acronyms and Definitions**

Define all acronyms or technical definitions used in this form.

ADD = automated dispensing device

### **Mandate and Impetus**

Identify the mandate for this regulatory change and any other impetus that specifically prompted its initiation (e.g., new or modified mandate, petition for rulemaking, periodic review, or board decision). For purposes of executive branch review, "mandate" has the same meaning as defined in Executive Order 14 (as amended, July 16, 2018), "a directive from the General Assembly, the federal government, or a court that requires that a regulation be promulgated, amended, or repealed in whole or part."

The impetus for the change is response to a petition for rulemaking by updating of regulations to facilitate new technologies in the practice of pharmacy. Since the technologies have already been approved for pilot programs in several hospital systems and have shown to be safe and effective, the Board's decision was to incorporate the allowances into regulation.

## Legal Basis

Identify (1) the promulgating agency, and (2) the state and/or federal legal authority for the regulatory change, including the most relevant citations to the Code of Virginia and Acts of Assembly chapter number(s), if applicable. Your citation must include a specific provision, if any, authorizing the promulgating agency to regulate this specific subject or program, as well as a reference to the agency's overall regulatory authority.

Regulations are promulgated under the general authority of Chapter 24 of Title 54.1 of the Code of Virginia. Section 54.1-2400, which provides the Board of Pharmacy the authority to promulgate regulations to administer the regulatory system:

#### § 54.1-2400 -General powers and duties of health regulatory boards

*The general powers and duties of health regulatory boards shall be:* 

6. To promulgate regulations in accordance with the Administrative Process Act (§ 2.2-4000 et seq.) that are reasonable and necessary to administer effectively the regulatory system, which shall include provisions for the satisfaction of board-required continuing education for individuals registered, certified, licensed, or issued a multistate licensure privilege by a health regulatory board through delivery of health care services, without compensation, to low-income individuals receiving health services through a local health department or a free clinic organized in whole or primarily for the delivery of those health services. Such regulations shall not conflict with the purposes and intent of this chapter or of Chapter 1 (§ 54.1-100 et seq.) and Chapter 25 (§ 54.1-2500 et seq.).

The specific authority for the Board to regulate the dispensing of prescription drugs is found in:

#### § 54.1-3307. Specific powers and duties of Board.

*A.* The Board shall regulate the practice of pharmacy and the manufacturing, dispensing, selling, distributing, processing, compounding, or disposal of drugs and devices. The Board shall also control the character and standard of all drugs, cosmetics and devices within the

Commonwealth, investigate all complaints as to the quality and strength of all drugs, cosmetics, and devices and take such action as may be necessary to prevent the manufacturing, dispensing, selling, distributing, processing, compounding and disposal of such drugs, cosmetics and devices that do not conform to the requirements of law.

The Board's regulations shall include criteria for:

1. Maintenance of the quality, quantity, integrity, safety and efficacy of drugs or devices distributed, dispensed or administered.

2. Compliance with the prescriber's instructions regarding the drug, its quantity, quality and directions for use.

3. Controls and safeguards against diversion of drugs or devices.

4. Maintenance of the integrity of, and public confidence in, the profession and improving the delivery of quality pharmaceutical services to the citizens of Virginia.

5. Maintenance of complete records of the nature, quantity or quality of drugs or substances distributed or dispensed, and of all transactions involving controlled substances or drugs or devices so as to provide adequate information to the patient, the practitioner or the Board.

6. Control of factors contributing to abuse of legitimately obtained drugs, devices, or controlled substances.

7. Promotion of scientific or technical advances in the practice of pharmacy and the manufacture and distribution of controlled drugs, devices or substances.

8. Impact on costs to the public and within the health care industry through the modification of mandatory practices and procedures not essential to meeting the criteria set out in subdivisions 1 through 7 of this section.

9. Such other factors as may be relevant to, and consistent with, the public health and safety and the cost of rendering pharmacy services.

*B.* The Board may collect and examine specimens of drugs, devices and cosmetics that are manufactured, distributed, stored or dispensed in the Commonwealth.

#### Purpose

Describe the specific reasons why the agency has determined that this regulation is essential to protect the health, safety, or welfare of citizens. In addition, explain any potential issues that may need to be addressed as the regulation is developed.

The purpose of this regulatory action is to update regulations for utilization of newer technologies in the practice of pharmacy in a hospital system and for facilitating time for pharmacists to be more involved in direct patient care. Two pilots for central distribution (involving more than a dozen facilities) have been approved by the Board and have been shown to protect the health and safety of the drug supply and patients in hospitals.

#### Substance

Briefly identify and explain the new substantive provisions that are being considered, the substantive changes to existing sections that are being considered, or both.

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Section 460 A currently requires a pharmacist to check all Schedule II-VI drugs delivered to a hospital unit as floor-stock before the drugs leave the pharmacy and to initial or sign manually or electronically the record of distribution verifying the accuracy of the distribution.

Section 490 C currently requires drugs in an ADD in a hospital to include the initials of the pharmacist at the hospital who checked the drugs removed from the pharmacy and the delivery record for accuracy.

Amendments to sections 460 and 490 would allow a pharmacist at a central distribution company to verify Schedule VI drugs to be placed in an ADD prior to delivery to the receiving hospital and pharmacy technicians at the hospital to load the drugs directly into the ADD without further verification by a pharmacist at the hospital.

### **Alternatives to Regulation**

Describe any viable alternatives to the regulatory change that were considered, and the rationale used by the agency to select the least burdensome or intrusive alternative that meets the essential purpose of the regulatory change. Also, include discussion of less intrusive or less costly alternatives for small businesses, as defined in § 2.2-4007.1 of the Code of Virginia, of achieving the purpose of the regulatory change.

The Board of Pharmacy received a petition for rulemaking on May of 2021 and published a request for comment until July 7, 2021. There were 40 comments posted on the Townhall; all were in favor of the petitioner's request as being a safer and more cost-effective delivery method. The proposal would constitute a less burdensome and intrusive alternative to the current regulation.

#### Periodic Review and Small Business Impact Review Announcement

This NOIRA is not being used to announce a periodic review or a small business impact review.

### **Public Participation**

Indicate how the public should contact the agency to submit comments on this regulation, and whether a public hearing will be held, by completing the text below. In addition, as required by § 2.2-4007.02 of the Code of Virginia describe any other means that will be used to identify and notify interested parties and seek their input, such as regulatory advisory panels or general notices.

The agency is seeking comments on this regulatory action, including but not limited to: ideas to be considered in the development of this proposal, the costs and benefits of the alternatives stated in this background document or other alternatives, and the potential impacts of the regulation.

The agency is also seeking information on impacts on small businesses as defined in § 2.2-4007.1 of the Code of Virginia. Information may include: projected reporting, recordkeeping, and other administrative costs; the probable effect of the regulation on affected small businesses;

and the description of less intrusive or costly alternatives for achieving the purpose of the regulation.

Anyone wishing to submit comments may do so via the Regulatory Town Hall website (<u>http://www.townhall.virginia.gov</u>) or by mail to Elaine Yeatts, 9960 Mayland Drive, Suite 300, Henrico, VA 23233; by email to <u>elaine.yeatts@dhp.virginia.gov</u>; by fax to (804) 527-4434. Written comments must include the name and address of the commenter. In order to be considered, comments must be received by midnight on the last day of the public comment period.

A public hearing will be held following the publication of the proposed stage of this regulatory action and notice of the hearing will be posted on the Virginia Regulatory Town Hall website (<u>http://www.townhall.virginia.gov</u>) and on the Commonwealth Calendar website (<u>https://www.virginia.gov/connect/commonwealth-calendar</u>). Both oral and written comments may be submitted at that time.

A Regulatory Advisory Panel will not be used for development of regulatory changes; the amendments will be drafted by the Regulation Committee.