19101805D HOUSE BILL NO. 2559 1 2 Offered January 9, 2019 3 Prefiled January 9, 2019 4 A BILL to amend and reenact §§ 54.1-3408.02, as it shall become effective, and 54.1-3410 of the Code 5 of Virginia, relating to electronic transmission of certain prescriptions; exceptions. 6 Patrons—Pillion and O'Quinn 7 8 Referred to Committee on Health, Welfare and Institutions 9 10 Be it enacted by the General Assembly of Virginia: 1. That §§ 54.1-3408.02, as it shall become effective, and 54.1-3410 of the Code of Virginia are 11 amended and reenacted as follows: 12 § 54.1-3408.02. (Effective July 1, 2020) Transmission of prescriptions. 13 14 A. Consistent with federal law and in accordance with regulations promulgated by the Board, 15 prescriptions may be transmitted to a pharmacy as an electronic prescription or by facsimile machine 16 and shall be treated as valid original prescriptions. B. Any prescription for a controlled substance that contains an opiate opioid shall be issued as an 17 18 electronic prescription. 19 C. The requirements of subsection B shall not apply if: 20 1. The prescriber dispenses the controlled substance that contains an opioid directly to the patient or 21 the patent's agent; 22 2. The prescription is for an individual who is residing in a hospital, assisted living facility, nursing 23 home, or residential health care facility or is receiving services from a hospice provider or outpatient 24 dialysis facility; 25 3. The prescriber experiences temporary technological or electrical failure or other temporary 26 extenuating circumstance that prevents the prescription from being transmitted electronically, provided 27 that the prescriber documents the reason for this exception in the patient's medical record; 28 4. The prescriber issues a prescription to be dispensed by a pharmacy located on federal property, 29 provided that the prescriber documents the reason for this exception in the patient's medical record; 30 5. The prescriber has issued fewer than 25 prescriptions during the most recent 12-month period 31 with a maximum of a seven-day supply for each prescription; 6. The prescription is issued by a licensed veterinarian for the treatment of an animal; 32 33 7. The FDA requires the prescription to contain elements that are not able to be included in an 34 *electronic* prescription: 35 8. The prescription is for an opioid under a research protocol; 36 9. The prescription is issued in accordance with an executive order of the Governor of a declared 37 emergency: 38 10. The prescription cannot be issued electronically in a timely manner and the patient's condition is 39 at risk, provided that the prescriber documents the reason for this exception in the patient's medical 40 record: or 41 11. The prescriber has been issued a waiver pursuant to subsection D. D. The licensing health regulatory board of a prescriber may grant such prescriber, in accordance 42 with regulations adopted by such board, a waiver of the requirements of subsection B, for a period not 43 to exceed one year, due to demonstrated economic hardship, technological limitations that are not 44 45 reasonably within the control of the prescriber, or other exceptional circumstances demonstrated by the 46 prescriber. 47 § 54.1-3410. When pharmacist may sell and dispense drugs. 48 A. A pharmacist, acting in good faith, may sell and dispense drugs and devices to any person 49 pursuant to a prescription of a prescriber as follows: 50 1. A drug listed in Schedule II shall be dispensed only upon receipt of a written prescription that is 51 properly executed, dated and signed by the person prescribing on the day when issued and bearing the 52 full name and address of the patient for whom, or of the owner of the animal for which, the drug is 53 dispensed, and the full name, address, and registry number under the federal laws of the person 54 prescribing, if he is required by those laws to be so registered. If the prescription is for an animal, it 55 shall state the species of animal for which the drug is prescribed; 2. In emergency situations, Schedule II drugs may be dispensed pursuant to an oral prescription in 56 57

3. Whenever a pharmacist dispenses any drug listed within Schedule II on a prescription issued by a 58

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accordance with the Board's regulations;

59 prescriber, he shall affix to the container in which such drug is dispensed, a label showing the prescription serial number or name of the drug; the date of initial filling; his name and address, or the 60 name and address of the pharmacy; the name of the patient or, if the patient is an animal, the name of 61 62 the owner of the animal and the species of the animal; the name of the prescriber by whom the 63 prescription was written, except for those drugs dispensed to a patient in a hospital pursuant to a chart 64 order; and such directions as may be stated on the prescription.

B. A drug controlled by Schedules III through VI or a device controlled by Schedule VI shall be 65 dispensed upon receipt of a written or oral prescription as follows: 66

1. If the prescription is written, it shall be properly executed, dated and signed by the person 67 prescribing on the day when issued and bear the full name and address of the patient for whom, or of 68 the owner of the animal for which, the drug is dispensed, and the full name and address of the person 69 70 prescribing. If the prescription is for an animal, it shall state the species of animal for which the drug is 71 prescribed.

72 2. If the prescription is oral, the prescriber shall furnish the pharmacist with the same information as 73 is required by law in the case of a written prescription for drugs and devices, except for the signature of 74 the prescriber.

75 A pharmacist who dispenses a Schedule III through VI drug or device shall label the drug or device 76 as required in subdivision A 3 of this section.

77 C. A drug controlled by Schedule VI may be refilled without authorization from the prescriber if, 78 after reasonable effort has been made to contact him, the pharmacist ascertains that he is not available 79 and the patient's health would be in imminent danger without the benefits of the drug. The refill shall be 80 made in compliance with the provisions of § 54.1-3411.

81 If the written or oral prescription is for a Schedule VI drug or device and does not contain the address or registry number of the prescriber, or the address of the patient, the pharmacist need not 82 83 reduce such information to writing if such information is readily retrievable within the pharmacy.

D. Pursuant to authorization of the prescriber, an agent of the prescriber on his behalf may orally 84 85 transmit a prescription for a drug classified in Schedules III through VI if, in such cases, the written record of the prescription required by this subsection specifies the full name of the agent of the 86 87 prescriber transmitting the prescription.

88 E. (Effective July 1, 2020) No pharmacist shall dispense a controlled substance that contains an 89 opiate unless the prescription for such controlled substance is issued as an electronic prescription. A 90 dispenser who receives a non-electronic prescription for a controlled substance containing an opioid is 91 not required to verify that one of the exceptions set forth in § 54.1-3408.02 applies and may dispense 92 such controlled substance pursuant to such prescription and applicable law.

2. That the Board of Medicine, the Board of Nursing, the Board of Dentistry, and the Board of 93 Optometry shall promulgate regulations to implement the provisions of this act regarding 94 95 prescriber waivers to be effective within 280 days of its enactment.

3. That the Secretary of Health and Human Resources shall convene a work group of interested 96 97 stakeholders, including the Medical Society of Virginia, the Virginia Hospital and Healthcare 98 Association, the Virginia Dental Association, the Virginia Association of Health Plans, and the 99 Virginia Pharmacists Association, to evaluate the implementation of the electronic prescription 100 requirement for controlled substances and shall report to the Chairmen of the House Committee 101 on Health, Welfare and Institutions and the Senate Committee on Education and Health by November 1, 2022. The work group's report shall identify the successes and challenges of 102 implementing the electronic prescription requirement and offer possible recommendations for 103

increasing the electronic prescribing of controlled substances that contain an opioid. 104