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## IN THE HOUSE OF REPRESENTATIVES

## HOUSE BILL NO. 191

## BY HEALTH AND WELFARE COMMITTEE

1	AN ACT
2	RELATING TO PHARMACY; AMENDING SECTION 54-1704, IDAHO CODE, TO PROVIDE THAT
3	PHARMACISTS MAY MAKE CERTAIN PRESCRIPTIONS AS AUTHORIZED BY RULE OF THE
4	BOARD OF PHARMACY.
5	Be It Enacted by the Legislature of the State of Idaho:
6	SECTION 1. That Section 54-1704, Idaho Code, be, and the same is hereby
7	amended to read as follows:
8	54-1704. PRACTICE OF PHARMACY. "Practice of pharmacy" means:
9	(1) The interpretation, evaluation and dispensing of prescription drug
10	orders;
11	(2) Participation in drug and device selection, drug administration,
12	prospective and retrospective drug reviews and drug or drug-related re-
13	search;
14	(3) The provision of patient counseling and the provision of those acts
15	or services necessary to provide pharmaceutical care;
16	(4) The responsibility for:
17	(a) Compounding and labeling of drugs and devices, except labeling by
18	a manufacturer, repackager or distributor of nonprescription drugs and
19	commercially packaged legend drugs and devices;
20	(b) Proper and safe storage of drugs and devices, and maintenance of
21	proper records for them; and
22	(c) The offering or performing of those acts, services, operations or
23	transactions necessary to the conduct, operation, management and con-
24	trol of pharmacy;
25	(5) The prescribing of:
26	(a) Dietary fluoride supplements when prescribed according to the Amer-
27	ican dental association's recommendations for persons whose drinking
28	water is proven to have a fluoride content below the United States de-
29	partment of health and human services' recommended concentration;
30	(b) Agents for active immunization when prescribed for susceptible per-
31	sons six (6) years of age or older for the protection from communicable
32	disease;
33	(c) Opioid antagonists pursuant to section 54-1733B, Idaho Code; and
34	(d) Epinephrine auto-injectors pursuant to sections 54-1733C and
35	54-1733D, Idaho Code; and
36	(e) Drugs, drug categories or devices that are specifically autho-
37	rized in rules adopted by the board. Such drugs and devices shall be
38	prescribed in accordance with the product's federal food and drug ad-
39	ministration-approved labeling. Drugs, drug categories or devices
40	authorized by the board under this section shall be limited to condi-

(i) Do not require a new diagnosis;

1	<pre>(ii) Are minor and generally self-limiting;</pre>
2	(iii) Have a test that is used to guide diagnosis or clinical deci-
3	sion-making and are waived under the federal clinical laboratory
4	improvement amendments of 1988; or
5	(iv) In the professional judgment of the pharmacist, threaten
6	the health or safety of the patient should the prescription not be
7	immediately dispensed. In such cases, only sufficient quantity
8	may be provided until the patient is able to be seen by another
9	provider.
10	The board shall not adopt any rules authorizing a pharmacist to pre-
11	scribe a controlled drug, compounded drug or biological product.