Sixty-third Legislative Assembly of North Dakota

SENATE BILL NO. 2190

Introduced by

Senators Dever, Berry, J. Lee

Representatives Damschen, Devlin, Rohr

- 1 A BILL for an Act to create and enact a new section to chapter 19-02.1 of the North Dakota
- 2 Century Code, relating to biosimilar biological products.

3 BE IT ENACTED BY THE LEGISLATIVE ASSEMBLY OF NORTH DAKOTA:

4 **SECTION 1.** A new section to chapter 19-02.1 of the North Dakota Century Code is created 5 and enacted as follows:

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6 Biosimilar biological products.

- 7 <u>1.</u> <u>In this section:</u>
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 a.
 "Biological product", "biosimilar", "interchangeable", "interchangeable biological

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 product", "license", and "reference product" mean the same as these terms mean

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 under section 351 of the Public Health Service Act [42 U.S.C. 262].
- 11b."Prescription" means a product that is subject to section 503(b) of the federal12Food, Drug, and Cosmetic Act [21 U.S.C. 353(b)].
- A pharmacy may substitute a prescription biosimilar product for a prescribed product
 only if:

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 a.
 The biosimilar product has been determined by the United States food and drug

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 administration to be interchangeable with the prescribed product for the specified

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 indicated use:

- <u>b.</u> The prescribing practitioner does not specifically indicate in the practitioner's own
 <u>handwriting "brand medically necessary" on a written prescription, does not</u>
 expressly indicate that an oral prescription is to be dispensed as communicated.
- 20 <u>expressly indicate that an oral prescription is to be dispensed as communicated.</u>
 21 <u>or has not taken a specific overt action to include the "brand medically</u>
- 22 necessary" language with an electronically transmitted prescription; and
- 23 <u>c.</u> <u>The pharmacist informs the individual receiving the biological product that the</u>
 24 <u>biological product may be substituted with a biosimilar product and that the</u>

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1	I	individual has a right to refuse the biosimilar product selected by the pharmacist
2		and the individual chooses not to refuse;
3	<u> <u>d.</u> </u>	The pharmacist notifies the prescribing practitioner in writing or via electronic
4		transmission within twenty-four hours of the substitution; and
5	<u> <u>e.</u> </u>	The pharmacy and the prescribing practitioner retain a written record of the
6		biosimilar substitution for a period of no less than five years.
7	<u>3.The bc</u>	pard of pharmacy shall:
8	<u>a.</u>	Maintain maintain on its public website a current list, or an internet link to a United
9	1	States food and drug administration-approved list, of biosimilar biological
10		products determined to be interchangeable under subdivision a of subsection 2;
11		and.
12	<u> </u>	Adopt rules for compliance, under which a pharmacy that violates subsection 2 is
13		subject to a specified civil money penalty.