## As Introduced

**135th General Assembly** 

Regular Session 2023-2024

H. B. No. 291

**Representatives Liston, Carruthers** 

Cosponsors: Representatives Galonski, McNally, Russo, Robb Blasdel, Baker, Troy, Brennan, Brown, Ray, Hillyer

## A BILL

To enact section 390	2.63 of the Revis	ed Code 1
regarding prescri	ption drugs and m	edication 2
switching.		3

## BE IT ENACTED BY THE GENERAL ASSEMBLY OF THE STATE OF OHIO:

Section 1. That section 3902.63 of the Revised Code be	
enacted to read as follows:	5
Sec. 3902.63. (A) As used in this section,	6
"interchangeable biological product" and "generically equivalent	7
drug" have the same meanings as in section 3715.01 of the	8
Revised Code.	9
(B) Notwithstanding section 3901.71 of the Revised Code,	10
with regard to health benefit plans amended, issued, or renewed	11
on or after the effective date of this section, a health plan	12
issuer shall not do any of the following during a plan year:	13
(1) Increase a covered person's burden of cost-sharing	14
with respect to a drug;	15
(2) Move a drug to a more restrictive tier of a health	16

benefit plan's formulary;	17
(3) Remove a drug from a health benefit plan's formulary	18
unless one of the following occurred:	19
(a) The United States food and drug administration issued	20
a statement about the drug calling into question the clinical	
safety of the drug.	22
(b) The drug manufacturer notified the United States food	23
and drug administration of a permanent discontinuance or	24
interruption of the manufacture of the drug as required by 21	25
<u>U.S.C. 356c.</u>	26
(c) The drug manufacturer has removed the drug from sale	27
in the United States.	28
(4) Limit or reduce coverage of a drug with respect to a	29
covered person in any other way, including subjecting it to a	30
prior authorization requirement.	31
(C) This section shall not be construed to do any of the	32
following:	33
(1) Prevent a health plan issuer from adding a drug to its	34
<pre>formulary;</pre>	35
(2) Prevent a health plan issuer from removing a drug from	36
its formulary if the drug manufacturer has removed the drug from	37
sale in the United States;	38
(3) Prevent a health care provider from prescribing	39
another drug covered by the health benefit plan that the	
provider considers medically appropriate for the covered person;	41
(4) In the case of a prescribed drug for which a	42
generically equivalent drug or interchangeable biological	43

product is available, prevent any of the following:	44	
(a) A pharmacist from substituting the generically	45	
equivalent drug or interchangeable biological product for the		
prescribed drug in accordance with section 4729.38 of the		
Revised Code;	48	
(b) A health plan issuer from requiring a covered person	49	
to use the generically equivalent drug or interchangeable		
biological product instead of the prescribed drug, even when the	51	
equivalent or product becomes available during a plan year;	52	
(c) A covered person from using the generically equivalent	53	
drug or interchangeable drug product instead of the prescribed		
drug, even when the equivalent or product becomes available		
during a plan year.		
(5) Prevent a pharmacist from substituting for a	57	
prescribed epinephrine autoinjector another epinephrine		
autoinjector pursuant to section 4729.382 of the Revised Code.	59	
(D) A violation of this section shall be considered an	60	
unfair and deceptive practice in the business of insurance for		
the purposes of section 3901.21 of the Revised Code.		