

1 State of Arkansas  
2 89th General Assembly  
3 Regular Session, 2013  
4

*As Engrossed: S2/4/13*

# A Bill

SENATE BILL 149

5 By: Senator Files  
6

## For An Act To Be Entitled

8 AN ACT TO REGULATE THE SUBSTITUTION OF BIOSIMILAR  
9 BIOLOGICAL PRODUCTS FOR CERTAIN PRESCRIBED PRODUCTS;  
10 AND FOR OTHER PURPOSES.  
11

## Subtitle

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13  
14 TO REGULATE THE SUBSTITUTION OF  
15 BIOSIMILAR BIOLOGICAL PRODUCTS FOR  
16 CERTAIN PRESCRIBED PRODUCTS.  
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18  
19 BE IT ENACTED BY THE GENERAL ASSEMBLY OF THE STATE OF ARKANSAS:  
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21 *SECTION 1. Arkansas Code Title 17, Chapter 92, is amended to add an*  
22 *additional subchapter to read as follows:*

23 *Subchapter 5 – Biosimilar Biological Products*  
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25 *17-92-507. Biosimilar biological products.*

26 *(a) As used in this section:*

27 *(1) "Biological product", "biosimilar", "interchangeable",*  
28 *"interchangeable biological product", and "reference product" have the*  
29 *meanings established under Section 351 of the Public Health Service Act, 42*  
30 *U.S.C. § 262; and*

31 *(2) "Prescription" means a product that is subject to Section*  
32 *503(b) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. § 353(b).*

33 *(b)(1) Except as provided in subsection (c) of this section, when a*  
34 *pharmacist receives a prescription for a biological product, the pharmacist*  
35 *may dispense a lower cost interchangeable biosimilar drug product.*

36 *(2) The total amount charged for the substituted interchangeable*



1 biosimilar product or for dispensing the prescribed product shall not exceed  
2 the amount normally and regularly charged under comparable circumstances by  
3 the pharmacist for that prescribed product or for the dispensing of the  
4 prescribed product.

5 (3) A pharmacist, a pharmacist's employee, or agent of a  
6 pharmacist shall notify the prescriber of the substitution of an  
7 interchangeable biosimilar product, including the full name and manufacturer,  
8 in writing or electronically not later than three (3) days after the date the  
9 product is dispensed.

10 (4) A pharmacist, the pharmacist's employee, or agent of a  
11 pharmacist, before dispensing an interchangeable biosimilar as a substitute  
12 for the prescribed biological product, shall inform the person for whom the  
13 medication is prescribed and the label of the dispensed shall appropriately  
14 indicate the substitution.

15 (5) A pharmacist shall record and retain for a period of two (2)  
16 years such records, the substitution of a reference product, including the  
17 full name and manufacturer of the prescribed product and of the  
18 interchangeable biosimilar product substituted for the prescribed product.

19 (c) A pharmacist shall not dispense an interchangeable biosimilar  
20 product under subsection (b) of this section:

21 (1) Unless the purchaser agrees to the total charge, if the  
22 total charge for the interchangeable biosimilar product exceeds the total  
23 charge of the prescribed product originally prescribed;

24 (2) For a prescription in writing signed by the prescriber, if  
25 the prescriber indicates in his or her own handwriting by name or initial  
26 that a substitution shall not be made;

27 (3) For a prescription other than one in writing signed by the  
28 prescriber, if the prescriber expressly indicates that the prescription is to  
29 be dispensed as communicated;

30 (4) If the individual for whom the reference product is  
31 prescribed indicates that the prescription shall be dispensed as written or  
32 communicated; or

33 (5) If the Arkansas State Board of Pharmacy has determined that  
34 the product shall not be substituted and has notified all pharmacists of that  
35 determination.

36 (d) The Arkansas State Board of Pharmacy shall:

