1	State of Arkansas	As Engrossed: \$2/4/13	
2	89th General Assembly	A Bill	
3	Regular Session, 2013		SENATE BILL 149
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5	By: Senator Files		
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7	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.		
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13	Subtitle		
14	TO RE	GULATE THE SUBSTITUTION OF	
15	BIOSI	MILAR BIOLOGICAL PRODUCTS FOR	
16	CERTA	IN PRESCRIBED PRODUCTS.	
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19	BE IT ENACTED BY THE GR	ENERAL ASSEMBLY OF THE STATE OF A	RKANSAS:
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21	SECTION 1. Arkan	nsas Code Title 17, Chapter 92, i	s amended to add an
22	additional subchapter to read as follows:		
23	<u>Subchapter</u>	5 — Biosimilar Biological Produc	<u>ets</u>
24			
25	<u>17-92-507. Bios</u>	imilar biological products.	
26	(a) As used in a	<u>this section:</u>	
27	<u>(1) "Biolo</u>	ogical product", "biosimilar", "i	nterchangeable",
28	"interchangeable biological product", and "reference product" have the		
29	meanings established under Section 351 of the Public Health Service Act, 42		
30	<u>U.S.C. § 262; and</u>		
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 co	<u>ost interchangeable biosimilar dr</u>	ug product.
36	(2) The to	otal amount charged for the subst	ituted interchangeable

As Engrossed: S2/4/13 SB149

1 biosimilar product or for dispensing the prescribed product shall not exceed

- 2 <u>the amount normally and regularly charged under comparable circumstances by</u>
- 3 <u>the pharmacist for that prescribed product or for the dispensing of the</u>
- 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 <u>(4) A pharmacist, the pharmacist's employee, or agent of a</u>
- 11 pharmacist, before dispensing an interchangeable biosimilar as a substitute
- 12 <u>for the prescribed biological product, shall inform the person for whom the</u>
- 13 <u>medication is prescribed and the label of the dispensed shall appropriately</u>
- 14 <u>indicate the substitution</u>.
- 15 <u>(5) A pharmacist shall record and retain for a period of two (2)</u>
- 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 <u>interchangeable biosimilar product substituted for the prescribed product.</u>
- 19 <u>(c) A pharmacist shall not dispense an interchangeable biosimilar</u>
- 20 product under subsection (b) of this section:
- 21 <u>(1) Unless the purchaser agrees to the total charge, if the</u>
- 22 <u>total charge for the interchangeable biosimilar product exceeds the total</u>
- 23 charge of the prescribed product originally prescribed;
- 24 (2) For a prescription in writing signed by the prescriber, if
- 25 <u>the prescriber indicates in his or her own handwriting by name or initial</u>
- 26 <u>that a substitution shall not be made;</u>
- 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 <u>prescribed indicates that the prescription shall be dispensed as written or</u>
- 32 communicated; or
- 33 (5) If the Arkansas State Board of Pharmacy has determined that
- 34 <u>the product shall not be substituted and has notified all pharmacists of that</u>
- 35 <u>determination</u>.
- 36 <u>(d) The Arkansas State Board of Pharmacy shall:</u>

As Engrossed: S2/4/13 SB149

1	(1)(A) Determine which biosimilar biological products are		
2	interchangeable.		
3	(B) The Arkansas State Board of Pharmacy shall make the		
4	determination under subdivision (d)(1)(A) of this section on the basis of th		
5	determination of the United States Food and Drug Administration regarding		
6	interchangeability with the prescribed biological product; and		
7	(2) Notify each licensed pharmacist and the Arkansas State		
8	Medical Board of the determination and any additions or deletions the		
9	Arkansas State Board of Pharmacy may make in its discretion.		
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