1	State of Arkansas	As Engrossed: H2/18/19	
2	92nd General Assembly	A Bill	
3	Regular Session, 2019		HOUSE BILL 1269
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5	By: Representative Magie		
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7		For An Act To Be Entitled	
8	AN ACT TO	ALLOW PHARMACISTS TO MAKE BIOLOGIC	AL
9	PRODUCT SU	BSTITUTIONS; AND FOR OTHER PURPOSE	S.
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12		Subtitle	
13	TO AI	LLOW PHARMACISTS TO MAKE BIOLOGICAL	
14	PRODU	JCT SUBSTITUTIONS.	
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17	BE IT ENACTED BY THE G	ENERAL ASSEMBLY OF THE STATE OF AR	KANSAS:
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19	SECTION 1. Arka	nsas Code § 17-92-101, concerning	the definitions
20	relating to pharmacists, pharmacies, and the practice of pharmacy, is amended		
21	to add new subdivision	s to read as follows:	
22	<u>(25) "Bio</u>	logical product" means a biologica	<u>l product as defined</u>
23	<u>by 42 U.S.C. 262(i)(1)</u>	, as existing on January 1, 2019;	and
24	<u>(26) "Int</u>	erchangeable biological product" m	<u>eans a biological</u>
25	product that is interc	hangeable as defined by 42 U.S.C.	262(i)(3), as
26	<u>existing on January l</u> ,	2019.	
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28	SECTION 2. Arka	nsas Code § 17-92-503 is amended t	o read as follows:
29	17-92-503. Gene	ric <u>drug product and biological pr</u>	oduct substitutions.
30	(a)(l) <u>(A)</u> Excep	t as provided in subsection (b) of	this section, when a
31	pharmacist receives a	prescription for a brand or trade	name drug product <u>or</u>
32	biological product, th	e pharmacist may dispense a lower	cost generically
33	equivalent drug produc	t <u>or interchangeable biological pr</u>	oduct only when there
34	<u>will be a cost savings</u>	for the patient.	
35	<u>(B)</u>	The pharmacist shall disclose the	amount of the cost
36	savings at the request	of the patient.	



1 (2) The total amount charged for the substituted generically 2 equivalent drug product or interchangeable biological product or for dispensing the drug product or biological product shall not exceed the amount 3 4 normally and regularly charged under comparable circumstances by the 5 pharmacist for that drug product or biological product or for the dispensing 6 of that drug product or biological product. 7 (3) A pharmacist may not dispense a drug product or 8 interchangeable biological product with a total charge that exceeds the total 9 charge of the drug product or biological product originally prescribed unless 10 agreed to by the purchaser. 11 (b) The pharmacist shall not dispense a generically equivalent drug 12 product or interchangeable biological product under subsection (a) of this 13 section if: 14 (1) The prescriber, in the case of a prescription in writing 15 signed by the prescriber, indicates in his or her own handwriting by name or 16 initial that no substitution shall be made; 17 (2) The prescriber, in the case of a prescription other than one 18 in writing signed by the prescriber, expressly indicates that the 19 prescription is to be dispensed as communicated; 20 (3) The person for whom the drug product or biological product 21 is prescribed indicates that the prescription is to be dispensed as written 22 or communicated; or 23 (4) The Arkansas State Board of Pharmacy has determined that the 24 drug product or biological product should not be substituted and has notified 25 all pharmacists of that determination. 26 (c)(1) The Arkansas State Board of Pharmacy shall determine which 27 drugs are generically equivalent and which biological products are 28 interchangeable biological products as defined in § 17-92-101, relying on 29 standards scientifically supported and generally accepted in the field of pharmacy, and shall notify each licensed pharmacist and the Arkansas State 30 31 Medical Board of this determination. 32 (2) In making this determination, the Arkansas State Board of 33 Pharmacy may use a nationally recognized reference source that meets the 34 requirements of this act, notifying each licensed pharmacist and the Arkansas 35 State Medical Board of the reference source to be used and any additions or 36 deletions the Arkansas State Board of Pharmacy may make in its discretion. 12-06-2016 09:33:41 JMB011

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1	(d)(l) Within five (5) business days after dispensing an		
2	interchangeable biological product that has been substituted for a biological		
3	product, the dispensing pharmacist or his or her designee shall record the		
4	specific interchangeable biological product provided to the patient,		
5	including without limitation the name of the interchangeable biological		
6	product and the manufacturer of the interchangeable biological product.		
7	(2) The record shall be electronically accessible to the		
8	prescriber through:		
9	(A) An interoperable electronic medical records system;		
10	(B) An electronic prescribing technology;		
11	(C) A pharmacy benefit management system; or		
12	(D) A pharmacy record.		
13	(3) If requested by a prescriber, a pharmacist shall communicate		
14	to the prescriber within five (5) business days using facsimile, telephone,		
15	electronic transmission, or other prevailing means that an interchangeable		
16	biological product has been dispensed.		
17	(4) A communication is not required when:		
18	(A) An interchangeable biological product does not exist		
19	for the prescribed biological product; or		
20	(B) A refill prescription for a biological product is not		
21	substituted with an interchangeable biological product on a subsequent		
22	filling of the prescription.		
23	(5) The pharmacist or pharmacy shall maintain a record of		
24	biological products dispensed for at least two (2) years.		
25	(6) Under subdivision (d)(2) of this section, the dispensing		
26	<u>pharmacist is not:</u>		
27	(A) Required to show proof that a prescriber has access to		
28	the record in any type of payment audit conducted by a payer or pharmacy		
29	<u>benefit manager; or</u>		
30	(B) Subject to disciplinary action or civil penalties for		
31	failure to ensure that the record is accessible or for failure to access the		
32	<u>record.</u>		
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34	SECTION 3. Arkansas Code § 17-92-505 is amended to read as follows:		
35	17-92-505. Labeling.		
36	(a)(l) The pharmacist filling a prescription for dispensing to an		

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1 ultimate patient may affix to the container a label showing: 2 (A) The pharmacy name, address, and telephone number; 3 (B) The date of dispensing; 4 The serial number of the prescription; (C) 5 (D) The name of the patient; 6 The name of the prescribing practitioner; (E) 7 (F) Either: 8 (i) The trade name of the medication drug product, 9 if any, or the generic name and identity of the manufacturer of the dispensed 10 medication drug product, if the medication drug product appears generically 11 listed on the drug formulary list as established by this subchapter; or 12 (ii) In the case of a biological product, the trade 13 name of the biological product, if any, or the proper name of the biological product and identity of the manufacturer of the dispensed biological product; 14 15 (G) The strength per unit dose of the medication; 16 (H) The quantity of the medication; and 17 (I) Directions for use. 18 (2) If a pharmacist dispenses a generically equivalent product 19 or interchangeable biological product, the person for whom the medication is 20 prescribed shall be informed prior to before dispensing or the label should 21 appropriately indicate the substitution. 22 (3) However, this subsection shall This subsection does not 23 apply to the dispensing of medication to inpatients in hospitals. 24 (4) Further, in an appropriate manner, In the case of dispensing 25 a drug product or biological product, the prescribing practitioner may 26 indicate that the name, manufacturer, and strength of the medication 27 dispensed shall be deleted from the label. 28 (b) Any authorized person filling a prescription An authorized person 29 who fills a prescription for dispensing to an ultimate patient shall affix to 30 the container a label showing: 31 (1) the The trade name of the medication or the generic name of 32 the medication unless directed to the contrary by the physician. Failure to 33 comply with this subsection shall be grounds for disciplinary action. 34 prescribing practitioner; or 35 (2) The trade name, if any, or the proper name of the biological 36 product unless directed to the contrary by the prescribing practitioner.

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2	SECTION 4. Arkansas Code § 17-92-506 is amended to read as follows:		
3	17-92-506. Price Available drug product and biological product lists.		
4	(a)(1) A pharmacist may display, within the confines of the pharmacy,		
5	lists of available drug products and biological products, other than		
6	controlled substances, and current charges for the drug products <u>or</u>		
7	biological products or for the dispensing of the drug products or biological		
8	products in specified quantities.		
9	(2) Upon request, a pharmacy may make such lists available to		
10	its customers and other members of the public.		
11	(b) The Arkansas State Board of Pharmacy shall maintain on the website		
12	of the board a link to the lists of all interchangeable biological products		
13	approved by the United States Food and Drug Administration.		
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15	/s/Magie		
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