Sixty-seventh Legislative Assembly of North Dakota

HOUSE BILL NO. 1032

Introduced by

Legislative Management

(Health Care Committee)

- 1 A BILL for an Act to create and enact a new chapter to title 19 of the North Dakota Century
- 2 Code, relating to prescription drug cost transparency; and to provide a penalty.

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

4 **SECTION 1.** A new chapter to title 19 of the North Dakota Century Code is created and

5 enacted as follows:

6 **Definitions**.

- 7 <u>As used in this chapter:</u>
- 8 <u>1.</u> "Board" means the state board of pharmacy.
- 9 <u>2.</u> <u>"Commissioner" means the insurance commissioner.</u>
- 10 <u>3.</u> <u>"Concession" includes a free good, delayed billing, and billing forgiveness.</u>
- 11 <u>4.</u> <u>"Drug" has the same meaning as provided under section 19-02.1-01.</u>
- 12 <u>5.</u> "Health care plan" means an individual, blanket, or group plan, policy, or contract for
 13 <u>health care services issued or delivered in this state by a health insurer.</u>
- 14 <u>6.</u> <u>"Health insurer" means an insurance company, nonprofit health service corporation,</u>
- 15 <u>health maintenance organization, third-party payer, health program administered by a</u>
- 16 <u>state agency, or other person engaged as principal in the business of insurance which</u>
- 17 <u>issues or delivers a health care plan in this state.</u>
- 18 7. "Manufacturer-packaged drug container" means a manufacturer-prepared supply of 19 medication packaged in a container with a unique product-identifying national drug
- 20 <u>code number.</u>
- 21 8. "Net spending" means the cost of drugs minus any discounts that lower the price of
- the drugs, including a rebate, fee, retained price protection, retail pharmacy network
 spread, and dispensing fee.

	•				
1	<u>9.</u>	"Pharmacy benefits manager" has the same meaning as provided under section			
2		<u>19-03.6-01.</u>			
3	<u>10.</u>	"Prescription drug" means a:			
4		a. Substance for which federal or state law requires a prescription before the			
5		substance may be legally dispensed to the public;			
6		b. Drug or device that under federal law is required, before being dispensed or			
7		delivered, to be labeled with the statement:			
8		(1) "Caution: federal law prohibits dispensing without prescription" or "Rx only"			
9		or other legend that complies with federal law; or			
10		(2) "Caution: federal law restricts this drug to use by or on the order of a			
11		licensed veterinarian"; or			
12		c. Drug or device required by federal or state law to be dispensed on prescription or			
13		restricted to use by a practitioner.			
14	<u>11.</u>	"Rebate" includes any discount, financial incentive, or concession that affects the price			
15		of a drug to a pharmacy benefits manager or health insurer for a drug manufactured			
16		by the pharmaceutical manufacturer.			
17	<u>12.</u>	"Specialty drug" has the same meaning as provided under section 19-02.1-16.2.			
18	<u>13.</u>	"Utilization management" means a set of formal techniques designed to monitor the			
19		use of, or evaluate the medical necessity, appropriateness, efficacy, or efficiency of,			
20		health care services, procedures, or settings.			
21	<u>14.</u>	"Wholesale acquisition cost" means, with respect to a prescription drug, the			
22		manufacturer's list price for the prescription drug to wholesalers or direct purchasers in			
23		the United States for the most recent month for which the information is available, as			
24		reported in wholesale price guides or other publications of drug pricing data, such as			
25		Medi-Span Price Rx, Gold Standard Drug Database, or First Databank drug data. The			
26		term does not include a rebate, prompt pay, or other discount or other reduction in			
27		price.			
28	Disc	isclosure of drug pricing information.			
29	<u>1.</u>	Each drug manufacturer shall submit a report to the board no later than the fifteenth			
30		day of January. April. July, and October with the current wholesale acquisition cost			

1		info	ormati	on for the United States food and drug administration-approved drugs sold in		
2		<u>or i</u>	or into the state by that manufacturer.			
3	<u>2.</u>	<u>a.</u>	<u>Not</u>	more than thirty days after an increase in wholesale acquisition cost of forty		
4			perc	cent or greater over the preceding five calendar years or ten percent or		
5			grea	ater in the preceding twelve months for a prescription drug with a wholesale		
6			<u>acq</u>	uisition cost of seventy dollars or more for a manufacturer-packaged drug		
7			<u>con</u>	tainer, a drug manufacturer shall submit a report to the board. The report		
8			mus	st contain the following information:		
9			<u>(1)</u>	Name of the drug:		
10			<u>(2)</u>	Whether the drug is a brand name or a generic;		
11			<u>(3)</u>	The effective date of the change in wholesale acquisition cost;		
12			<u>(4)</u>	Aggregate, company-level research and development costs for the previous		
13				calendar year;		
14			<u>(5)</u>	Aggregate rebate amounts paid to each pharmacy benefits manager for the		
15				<u>calendar year;</u>		
16			<u>(6)</u>	The name of each of the manufacturer's drugs approved by the United		
17				States food and drug administration in the previous five calendar years;		
18			<u>(7)</u>	The name of each of the manufacturer's drugs that lost patent exclusivity in		
19				the United States in the previous five calendar years; and		
20			<u>(8)</u>	A statement of rationale regarding the factor or factors that caused the		
21				increase in the wholesale acquisition cost, such as raw ingredient shortage		
22				or increase in pharmacy benefits manager rebates.		
23		<u>b.</u>	<u>The</u>	quality and types of information and data a drug manufacturer submits to the		
24			<u>boa</u>	rd pursuant to this subsection must be the same as the quality and types of		
25			info	rmation and data the manufacturer includes in the manufacturer's annual		
26			<u>con</u>	solidated report on securities and exchange commission form 10-K or any		
27			othe	er public disclosure.		
28	<u>3.</u>	<u>A d</u>	rug m	anufacturer shall notify the board in writing if the manufacturer is introducing		
29		<u>a n</u>	ew pr	escription drug to market at a wholesale acquisition cost that exceeds the		
30		<u>thre</u>	esholo	set for a specialty drug under the Medicare part D program.		

1		<u>a.</u>	The notice must include a statement of rationale regarding the factor or factors
2			that caused the new drug to exceed the Medicare part D program price.
3		<u>b.</u>	The drug manufacturer shall provide the written notice within three calendar days
4			following the release of the drug in the commercial market.
5		<u>C.</u>	A drug manufacturer may make the notification pending approval by the United
6			States food and drug administration if commercial availability is expected within
7			three calendar days following the approval.
8	<u>4.</u>	<u>Wit</u>	hin thirty days of receipt of a report under this section, the board shall provide the
9		rep	orted information to the commissioner in a format ready for publication on the
10		<u>cor</u>	nmissioner's website.
11	<u>Dis</u>	closi	ure of pharmacy benefits manager information.
12	<u>1.</u>	<u>On</u>	or before April first of each year, a pharmacy benefits manager providing services
13		for	a health care plan shall file a report with the board. The report must contain the
14		<u>foll</u>	owing information for the previous calendar year:
15		<u>a.</u>	The aggregated rebates, fees, price protection payments, and any other
16			payments collected from each drug manufacturer;
17		<u>b.</u>	The aggregated dollar amount of rebates, price protection payments, fees, and
18			any other payments collected from each drug manufacturer which were passed
19			to health insurers;
20		<u>C.</u>	The aggregated fees, price concessions, penalties, effective rates, and any other
21			financial incentive collected from pharmacies which were passed to enrollees at
22			the point of sale;
23		<u>d.</u>	The aggregated dollar amount of rebates, price protection payments, fees, and
24			any other payments collected from drug manufacturers which were retained as
25			revenue by the pharmacy benefits manager; and
26		<u>e.</u>	The aggregated rebates passed on to employers.
27	<u>2.</u>	Re	ports submitted by pharmacy benefits managers under this section may not
28		<u>dis</u>	close the identity of a specific health benefit plan or enrollee, the prices charged for
29		<u>spe</u>	cific drugs or classes of drugs, or the amount of any rebates or fees provided for
30		<u>spe</u>	cific drugs or classes of drugs.

	•		
1	<u>3.</u>		nirty days of receipt of a report under this section, the board shall provide the
2		<u>reported</u>	information to the commissioner in a format ready for publication on the
3		<u>commise</u>	sioner's website. The information the board provides to the commissioner may
4		<u>not discl</u>	ose or tend to disclose proprietary or confidential information of any pharmacy
5		<u>benefit n</u>	nanager.
6	<u>Dis</u>	<u>closure o</u>	f health insurer spending information.
7	<u>1.</u>	<u>a. On</u>	or before April first of each year, each health insurer shall submit a report to
8		<u>the</u>	board. The report must contain the following information for the previous two
9		cale	endar years:
10		<u>(1)</u>	Names of the twenty-five most frequently prescribed drugs across all plans;
11		<u>(2)</u>	Names of the twenty-five prescription drugs dispensed with the highest
12			dollar spend in terms of gross revenue;
13		<u>(3)</u>	Percent increase in annual net spending for prescription drugs across all
14			plans:
15		<u>(4)</u>	Percent increase in premiums which is attributable to prescription drugs
16			across all plans;
17		<u>(5)</u>	Percentage of specialty drugs with utilization management requirements
18			across all plans; and
19		<u>(6)</u>	Premium reductions attributable to specialty drug utilization management.
20		<u>b. Wit</u>	hin thirty days of receipt of a report under this section, the board shall provide
21		<u>the</u>	reported information to the commissioner in a format ready for publication on
22		the	commissioner's website. The combined aggregated data from the reports
23		<u>whi</u>	ch the board provides to the commissioner must be provided in a manner that
24		doe	es not disclose or tend to disclose proprietary or confidential information of any
25		hea	alth insurer.
26	<u>2.</u>	<u>A report</u>	submitted by a health insurer may not disclose the identity of a specific health
27		<u>benefit p</u>	plan or the prices charged for specific prescription drugs or classes of
28		prescript	tion drugs.
29	We	<u>bsite.</u>	
30	<u>1.</u>	The com	missioner shall develop a website to publish information the board reports to
31		the com	missioner under this chapter. The commissioner shall make the website

1		available on the commissioner's website with a dedicated link prominently displayed		
2		on the home page, or by a separate, easily identifiable internet address.		
3	<u>2.</u>	Within thirty days of receipt of reported information from the board, the commissioner		
4		shall publish the reported information on the website developed under this section.		
5	Rulemaking - Forms - Services - Records.			
6	<u>1.</u>	The board and the commissioner may adopt rules to implement this chapter.		
7	<u>2.</u>	In consultation with the commissioner, the board shall develop forms that must be		
8		used for reporting required under this chapter.		
9	<u>3.</u>	The board may contract for services to implement this chapter.		
10	<u>4.</u>	A report received by the board is an exempt record as defined by section 44-04-17.1.		
11	Civil penalty.			
12	A health care plan, drug manufacturer, or pharmacy benefits manager that violates this			
13	chapter is subject to the imposition by the attorney general of a civil penalty not to exceed			
14	ten thousand dollars for each violation. The fine may be collected and recovered in an action			
15	brought in the name of the state.			