GENERAL ASSEMBLY OF NORTH CAROLINA **SESSION 2023**

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HOUSE BILL DRH10131-MG-48B

Short Title:	Educate Patients About Opioid Antagonists.	(Public)
Sponsors:	Representative Sasser.	
Referred to:		

1		A BILL TO BE ENTITLED				
2	AN ACT	REQUIRING HEALTH CARE PRACTITIONERS AND PHARMACISTS TO				
3	EDUC	CATE PATIENTS WITH PRESCRIPTIONS FOR OPIOID PAIN MEDICATIONS				
4	AND	MEDICATIONS TO TREAT OPIOID USE DISORDER ABOUT THE POTENTIAL				
5	DANO	DANGERS OF OPIOIDS, OVERDOSE PREVENTION, AND THE AVAILABILITY				
6	AND	USE OF OPIOID ANTAGONISTS TO PREVENT OVERDOSE DEATHS.				
7		eral Assembly of North Carolina enacts:				
8	SECTION 1. Article 1 of Chapter 90 of the General Statutes is amended by adding					
9	a new sect	tion to re	ad:			
10	" <u>§ 90-12.8</u>	8. Requ	rement to provide opioid antagonist education.			
11	<u>(a)</u>	Consist	ent with the federal Food and Drug Administration's labeling requirements for			
12	opioid pain medication and medication to treat opioid use disorder announced in its Drug Safety					
13	Communication dated July 23, 2020, a practitioner as defined in G.S. 90-87(22) shall do all of					
14	the follow	ing whe	n issuing a prescription for a Schedule II controlled substance described in			
15	<u>G.S. 90-9</u>	0 <u>(1):</u>				
16		<u>(1)</u>	Provide information regarding all of the following to each patient receiving			
17			the prescription:			
18			a. <u>The potential dangers of opioids.</u>			
19			b. <u>Overdose prevention.</u>			
20			c. The availability and use of a drug approved by the federal Food and			
21			Drug Administration as an opioid antagonist for the complete or partial			
22			reversal of opioid-induced respiratory depression.			
23		<u>(2)</u>	Provide the information described in sub-subdivisions (1)a. through (1)c. of			
24			this subsection to one or more persons if designated by the patient receiving			
25			the prescription or, for a patient who is a minor, to the minor's parent,			
26			guardian, or person standing in loco parentis.			
27	<u>(b)</u>	When	dispensing a Schedule II controlled substance described in G.S. 90-90(1), a			
28	pharmacy	, through	a pharmacist or pharmacy personnel, shall do one of the following:			
29		<u>(1)</u>	Make available the information described in sub-subdivisions (a)(1)a. through			
30			(a)(1)c. of this section that is consistent with the federal Food and Drug			
31			Administration's labeling requirements for opioid pain medication and			
32			medication to treat opioid use disorder announced in its Drug Safety			
33			Communication dated July 23, 2020.			
34		<u>(2)</u>	Post signage in a conspicuous place containing the information described in			
35			sub-subdivisions (a)(1)a. through (a)(1)c. of this section.			
36	<u>(c)</u>	<u>Nothin</u>	g in this section shall be construed to do any of the following:			



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1	<u>(1)</u>	Limit a practitioner's liability for negligent diagnosis or treatm	nent of a patient,
2		as allowed under applicable State or federal law.	
3	<u>(2)</u>	Constitute negligence per se or create a private right of act	tion against any
4		practitioner, including a pharmacy, a pharmacist, or pharmacy	personnel, who
5		fails to follow the requirements of this section."	
6	SECT	TON 2. This act becomes effective October 1, 2023.	