GENERAL ASSEMBLY OF NORTH CAROLINA SESSION 2025

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HOUSE BILL 592 PROPOSED COMMITTEE SUBSTITUTE H592-PCS40524-DC-11

Short Title: Toxic-Free Medical Devices Act of 2025.

(Public)

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Sponsors:

Referred to:

April 1, 2025 1 A BILL TO BE ENTITLED 2 AN ACT TO PROHIBIT THE MANUFACTURING, SELLING, AND DISTRIBUTING OF 3 INTRAVENOUS SOLUTION CONTAINERS AND INTRAVENOUS TUBING THAT 4 ARE INTENTIONALLY MADE WITH DEHP. 5 The General Assembly of North Carolina enacts: 6 SECTION 1. Chapter 130A of the General Statutes is amended by adding a new 7 Article to read as follows: 8 "Article 19C. 9 "DEHP Hazard Management. 10 "§ 130A-453.33. Legislative finding. 11 The General Assembly finds all of the following: DEHP and other ortho-phthalates are toxic chemicals used primarily to 12 (1)produce flexibility in plastics, mainly polyvinyl chloride (PVC). 13 14 (2)DEHP is the most common plasticizer used in medical devices, including 15 intravenous solution containers, which are also known as IV bags, and intravenous tubing. 16 Over the course of its shelf life, DEHP leaches from IV bags and tubing made 17 (3) 18 from DEHP into the solutions being held in the medical devices. 19 DEHP is classified by the United States Environmental Protection Agency as (4) 20 an endocrine-disrupting compound since it can: Interfere with the hormonal system in humans and animals. 21 a. Mimic or block the actions of hormones, leading to adverse effects on 22 b. 23 reproductive health, development, and metabolism. 24 DEHP exposure has been associated with adverse effects on reproductive (5) 25 organs and fertility. DEHP can also disrupt normal reproductive development, reduce sperm quality, and affect hormone levels in both males and females. 26 27 (6) DEHP is metabolized in the liver and can accumulate in the body over time. Prolonged exposure to high levels of DEHP has been shown to cause liver and 28 kidney damage in animal studies. 29 30 (7)Inhalation or ingestion of DEHP can cause respiratory irritation and allergic 31 reactions in some individuals, particularly those with preexisting respiratory 32 conditions or sensitivities. 33 (8) Studies have suggested a potential link between DEHP exposure and certain 34 types of cancer, including breast, liver, lung, and testicular cancer. The United States Environmental Protection Agency has determined that 35 (9) 36 DEHP is a probable human carcinogen.



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1	<u>(10)</u>	The leaching of DEHP from medical devices at varying	ng concentrations has
2		been linked to multidrug resistance in breast cancer	cells, inhibiting the
3		effectiveness of breast cancer drugs. This phenomenon	has been observed at
1		both high and low concentrations of DEHP, highlightin	g the potential impact
5		of DEHP leaching on cancer treatment outcomes.	
5	<u>(11)</u>	Exposure to DEHP has been linked to multidrug resista	
7		breast cancer cells, inhibiting the apoptosis mechanis	
3		cancer drugs, such as tamoxifen, and increasing cell pro-	
9	<u>(12)</u>	DEHP has been suggested to serve as a mitogenic	
0		receptor-positive breast cancer cells, potentially mal	king them multidrug
1		resistant.	
2	" <u>§ 130A-453.34.</u>		
3		g definitions apply in this Article:	
4	<u>(1)</u>	DEHP. – Di(2-ethylhexyl) phthalate.	
5	<u>(2)</u>	Health care practitioner An individual who is author	
6		component of the healing arts by a license, permit, cert	ificate, or registration
7		issued by a State licensing agency or board.	
8	<u>(3)</u>	Intentionally added DEHP. – DEHP that a manufactu	
9	(4)	added to a product and that has a functional or technical	
0	<u>(4)</u>	Intravenous solution container. – A container used to h	
1 2		or nutrition therapy that is intravenously delivered to a	patient in a nospital,
	(5)	outpatient facility, or other health care facility.	
3 4	<u>(5)</u>	Intravenous tubing. – Tubing used to intravenous	•
4 5	(6)	<u>medication, or nutrients directly to an adult, child, or in</u> <u>Ortho-phthalate. – A class of chemicals that are esters of</u>	
5 6	<u>(6)</u>	including DEHP or any of the following:	<u>n ormo-philiane aciu,</u>
7		<u>a. Benzyl butyl phthalate (BBP).</u>	
8		<u>b.</u> <u>Dibutyl phthalate (DBP).</u>	
9			
0		<u>c.</u> <u>Dicyclohexyl phthalate (DCHP).</u> <u>d.</u> <u>Diethyl phthalate (DEP).</u>	
1			
2		e.Diisobutyl phthalate (DIBP).f.Diisodecyl phthalate (DIDP).	
3			
4		h. Di-n-hexyl phthalate (DnHP).	
5		g.Diisononyl phthalate (DINP).h.Di-n-hexyl phthalate (DnHP).i.Di-n-octyl phthalate (DNOP).j.Di-n-pentyl phthalate (DnPP).	
6		<u>j.</u> <u>Di-n-pentyl phthalate (DnPP).</u>	
7		k. Diisoheptyl phthalate (DIHP).	
8	(7)	Unintentionally added DEHP. – DEHP in an intraveno	ous solution container
9		or intravenous tubing product that is not used for function	nal or technical effect
0		on the product.	
1	" <u>§ 130A-453.35.</u>	Prohibitions.	
-2	<u>(a)</u> Intrav	enous Solution Containers Beginning January 1, 2030, a	a person or entity shall
13		sell, or distribute into commerce in the State of North	Carolina intravenous
4		rs made with intentionally added DEHP.	
5		enous Tubing. – Beginning January 1, 2035, a perso	-
46		, or distribute into commerce in the State of North Caroli	na intravenous tubing
17 10		ionally added DEHP.	
18 10		<u>cement. – A person may not replace DEHP, pursuant</u>	to this Article, with
19	another ortho-ph	thalate in a new or revised medical device.	

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1	(d) Maximum Quantity. – An intravenous solution container or intravenous tubir	ng		
2	product shall not have unintentionally added DEHP present at a quantity at or above 0.1 perce	nt		
3	weight per weight (w/w).			
4	(e) <u>Exemptions. – The following items, as described in Title 21 of the Code of Feder</u>	al		
5	Regulations, are exempt from these provisions:			
6	(1) <u>Human blood collection and storage bags.</u>			
7	(2) <u>Apheresis and cell therapy blood kits and bags, including integral tubing.</u>			
8	(f) Delayed Compliance. – A person or entity, due to pending United States Food ar			
9	Drug Administration approval for the DEHP-free intravenous solution container or due to the			
10	manufacturer not having adequate equipment to manufacture the DEHP-free intravenous solution			
11	container, shall meet the requirement in subsection (a) of this section by January 1, 2032, if a	all		
12	of the following conditions are met:			
13	(1) The person or entity notified its North Carolina customers, no later that			
14	October 1, 2025, that it has commenced development of the DEHP-free	<u>ee</u>		
15	intravenous solution container to meet the requirements of this section.			
16	(2) The person or entity provides notice to its customers and posts to its offici			
17	internet website, no later than January 1, 2028, that it will not meet the	ne		
18	deadline imposed pursuant to subsection (a) of this section."			
19	SECTION 2. G.S. 130A-22(b3) reads as rewritten:	1		
20	"(b3) The Secretary may impose an administrative penalty on a person who violates Artic			
21	19A or 19B Article 19A, 19B, or 19C of this Chapter or any rules adopted pursuant to Artic			
22 23	19A or 19B Article 19A, 19B, or 19C of this Chapter. Each day of a continuing violation is			
23 24	separate violation. The penalty shall not exceed five thousand dollars (\$5,000) for each day the			
24 25	violation continues for Article 19A of this Chapter. The penalty shall not exceed five thousar dollars (\$5,000) for each day the violation continues for Article 19B of this Chapter. <u>The penal</u>			
23 26	shall not exceed five thousand dollars (\$5,000) for each day the violation continues for Artic			
20 27	19C of this Chapter. The penalty authorized by this section does not apply to a person who is n			
28	required to be certified under Article 19A or 19B."	.01		
28 29	SECTION 3. Except as otherwise provided, this act is effective when it become	es		
30	law.	05		
50	14 11 ·			