## GENERAL ASSEMBLY OF NORTH CAROLINA

## SESSION 1997

S 1 SENATE BILL 945 Short Title: Prescription Refill Safety Act. (Public) Sponsors: Senators Rand; Ballantine, Cooper, and Forrester. Referred to: Commerce. April 17, 1997 A BILL TO BE ENTITLED AN ACT TO REQUIRE THE PRESCRIBER'S AND THE PATIENT'S CONSENT FOR INTERCHANGE OF A LIMITED CLASS OF DRUGS. The General Assembly of North Carolina enacts: Section 1. G.S. 90-85.27 reads as rewritten: "§ 90-85.27. Definitions. As used in G.S. 90-85.28 through G.S. 90-85.31: 'Equivalent drug product' means a drug product which has the same (1) established name, active ingredient, strength, quantity, and dosage form, and which is therapeutically equivalent to the drug product identified in the prescription; 'Established name' has the meaning given in section 502(e)(3) of the (2) Federal Food, Drug and Cosmetic Act, 21 U.S.C. 352(e)(3); 'Good manufacturing practice' has the meaning given it in Part 211 of (3) Chapter 1 of Title 21 of the Code of Federal Regulations: 'Manufacturer' means the actual manufacturer of the finished dosage (4) form of the drug; 'Narrow therapeutic index drugs' means those pharmaceuticals having a (4a) narrowly defined range between risk and benefit. Such drugs have less than a two-fold difference in the minimum toxic concentration and

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	minimum effective concentration in the blood or are those drug product
	formulations that exhibit limited or erratic absorption, formulation-
	dependent bioavailability, and wide intrapatient pharmacokinetic
	variability that requires blood-level monitoring. Drugs identified as
	having narrow therapeutic indices shall be designated by the North
	Carolina Secretary of Human Resources upon the advice of the State
	Health Director, North Carolina Board of Pharmacy, and North Carolina
	Medical Board, as narrow therapeutic index drugs and shall be subject
	to the provisions of G.S. 90-85.28(b1). The list of narrow therapeutic
	index drugs shall be published annually in January in the North Carolina
	Register;
(5)	'Prescriber' means anyone authorized to prescribe drugs pursuant to the
	(5)

laws of this State."

Section 2. G.S. 90-85.28 is amended by adding the following new subsection to read:

"(b1) A prescription for a narrow therapeutic index drug shall be refilled using only the same drug product by the same manufacturer that the pharmacist last dispensed under the prescription, unless the prescriber is notified by the pharmacist prior to the dispensing of another manufacturer's product, and the prescriber and the patient give written consent to the dispensing of the other manufacturer's product. For purposes of this subsection, the term 'refilled' shall include a new prescription written at the expiration of a prescription which continues the patient's therapy on a narrow therapeutic index drug."

Section 3. This act becomes effective July 1, 1997.

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