1997 SESSION

	973712200
1	HOUSE BILL NO. 2736
2	House Amendments in [] — February 3, 1997
3 4 5	A BILL to amend and reenact §§ 32.1-79 and 32.1-87 of the Code of Virginia, relating to the Voluntary Formulary.
5 6 7	Patrons—DeBoer, Darner and Davies; Senators: Couric, Gartlan, Houck and Lambert
, 8 9	Referred to Committee on Health, Welfare and Institutions
9 10	Be it enacted by the General Assembly of Virginia:
11	1. That §§ 32.1-79 and 32.1-87 of the Code of Virginia is amended and reenacted as follows:
12	§ 32.1-79. Definitions.
13	As used in this article:
14	"Formulary Board" means the Virginia Voluntary Formulary Board.
15	"Drug" shall have the same meaning as provided in § 54.1-3401.
16 17	"Drug product" means a specific drug in dosage form from a known source of manufacture, whether by brand or established name.
18	"Formulary" and "Voluntary Formulary" mean the Virginia Voluntary Formulary prepared in
19	accordance with the provisions of this article.
20	"Narrow therapeutic index drugs" means those pharmaceuticals having a more defined range
21	between risk and benefit. Such drugs are known to have less than a two-fold difference in the minimum
22	toxic concentration and minimum effective concentration in the blood.
23 24	"Therapeutic index of a drug" means the range between benefit and risk of the drug. § 32.1-87. Use of Formulary.
25	A. Use of the Voluntary Formulary by professional and institutional providers of health care shall be
26	voluntary. The prescription form shall include two boxes, one labelled "Voluntary Formulary Permitted"
27	and the other labelled "Dispense As Written." A prescriber may indicate his permission for the
28	dispensing of a drug product included in the Formulary upon signing a prescription form and marking
29 30	the box labelled "Voluntary Formulary Permitted." A Voluntary Formulary product shall be dispensed if the prescriber fails to indicate his preference. If no Voluntary Formulary product is immediately
30 31	available, or if the patient objects to the dispensing of a generic drug, the pharmacist may dispense a
32	brand name drug.
33	On and after July 1, 1993, printed prescription forms shall provide:
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35 36	" 🗌 Dispense As Written
30 37	□ Voluntary Formulary Permitted
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41	Signature of prescriber
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44 45	If neither box is marked, a Voluntary Formulary product must be dispensed."
45 46	If a prescriber orders a drug listed in the Formulary by its generic name, the pharmacist shall dispense a drug product from among those listed in the Formulary.
47	In the case of an oral prescription, the prescriber's oral dispensing instructions shall be followed. If
48	the pharmacist dispenses a drug product other than the brand name prescribed, he shall so apprise the
49	purchaser and shall indicate, unless otherwise directed by the prescriber, on both his permanent record
50	and the prescription label the brand name or, in the case of a generic drug product, the name of the
51 52	manufacturer or distributor. B. When a pharmacist dispenses a drug product other than the drug product prescribed under the
52 53	provisions of subsection A hereof pursuant to the Formulary, the drug product dispensed shall be at a
53 54	lower retail price than that of the drug product prescribed. Such retail price shall not exceed the usual
55	and customary retail price charged by the pharmacist for such generic or equivalent drug product
56	dispensed.
57	The selection of such drug product shall be the responsibility of the pharmacist dispensing the

57 The selection of such drug product shall be the responsibility of the pharmacist dispensing the 58 product and no employer, agent or other person may require the dispensing of a particular drug product HB2736E

59 which in the professional judgment of the dispensing pharmacist is not in the best interest of the patient. 60 C. Narrow therapeutic index drugs include: [carbamazephine carbamazepine], [conjugated

61 estrogens,] digoxin, levothyroxine, phenytoin, theophylline sustained release, and [Warfarin warfarin]

. A pharmacist shall not substitute or interchange a narrow therapeutic index drug, as defined in 62

63 § 32.1-79 and identified in this subsection, without the [written informed consent of the patient's 64

physician documented consent of the patient's prescriber] to the substitution or interchange [, prior to dispensing, in accordance with regulations to be promulgated by the Board of Pharmacy]. [The 65

written informed consent shall be signed by the patient's physician. 66

Other drugs identified hereinafter by the Food and Drug Administration as having narrow 67 therapeutic indices may be designated by the Voluntary Formulary Board as such narrow therapeutic 68

index drugs and shall be subject to the provisions of this subsection. 69

70 [2. That the provisions of this act shall become effective on July 1, 1998.]