

Bill No. CS/HB 1467, 2nd Eng.

Amendment No.

	<u>Senate</u>	CHAMBER ACTION	<u>House</u>
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11	Senator Brown-Waite moved the following amendment:		
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13	Senate Amendment (with title amendment)		
14	On page 185, between lines 28 and 29,		
15			
16	insert:		
17	Section 127. Subsection (6) of section 465.025,		
18	Florida Statutes, is amended to read:		
19	465.025 Substitution of drugs.--		
20	(6) The Board of Pharmacy and the Board of Medicine		
21	shall establish by rule a formulary of generic drug type and		
22	brand name drug products which are determined by the boards to		
23	demonstrate clinically significant biological or therapeutic		
24	inequivalence and which, if substituted, would pose a threat		
25	to the health and safety of patients receiving prescription		
26	medication.		
27	(a) The formulary may be added to or deleted from as		
28	the Board of Pharmacy and the Board of Medicine deem		
29	appropriate. Any person who requests any inclusion, addition,		
30	or deletion of a generic drug type or brand name drug product		
31	to the formulary shall have the burden of proof to show cause		

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1 why such inclusion, addition, or deletion should be made.

2 (b) Upon adoption of the formulary required by this
3 subsection, and upon each addition, deletion, or modification
4 to the formulary, the Board of Pharmacy shall mail a copy to
5 each manager of the prescription department of each community
6 pharmacy licensed by the state, each nonresident pharmacy
7 registered in the state, and each board regulating
8 practitioners licensed by the laws of the state to prescribe
9 drugs shall incorporate such formulary into its rules. No
10 pharmacist shall substitute a generically equivalent drug
11 product for a prescribed brand name drug product if the brand
12 name drug product or the generic drug type drug product is
13 included in the said formulary.

14 (c) The Department of Health shall study the necessity
15 of retaining drugs currently listed on the negative formulary,
16 taking into consideration whether each such drug has been
17 listed by the Federal Food and Drug Administration in the
18 Orange Book as an Approved Drug Product with Therapeutic
19 Equivalence Evaluations. The Department of Health shall report
20 its conclusions and recommendations to the Legislature by
21 November 1, 1999.

22
23 (Redesignate subsequent sections.)

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25
26 ===== T I T L E A M E N D M E N T =====

27 And the title is amended as follows:

28 On page 1, line 3, delete the word "practitioners;"

29
30 insert:

31 ; amending s. 465.025, F.S.; requiring the

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1 Department of Health to study the necessity of
2 retaining certain drugs on the negative
3 formulary; requiring a report to the
4 Legislature;
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