



General Assembly

February Session, 2022

Raised Bill No. 188

LCO No. 1654



Referred to Committee on GENERAL LAW

Introduced by:
(GL)

***AN ACT REQUIRING BRAND NAME PRESCRIPTION DRUG
MANUFACTURERS TO PROVIDE SAMPLES OF BRAND NAME
DRUGS TO GENERIC PRESCRIPTION DRUG MANUFACTURERS.***

Be it enacted by the Senate and House of Representatives in General Assembly convened:

1 Section 1. (NEW) (*Effective October 1, 2022*) (a) As used in this section:

2 (1) "Eligible product developer" means a person who seeks to develop
3 an application for the approval of a drug under subsections (b) and (j)
4 of Section 505 of the federal Food, Drug and Cosmetic Act or the
5 licensing of a biological product under Section 351 of the federal Public
6 Health Service Act; and

7 (2) "Wholesale acquisition cost" means the manufacturer's list price
8 for a brand-name drug or a generic drug per person, year or course of
9 treatment, when sold to wholesalers or direct purchasers in the United
10 States, not including discounts or rebates, for the most recent month for
11 which information is available.

12 (b) A manufacturer or wholesaler registered under chapter 417 of the
13 general statutes shall make a drug manufactured or developed by such

14 manufacturer or wholesaler and distributed in this state available for
15 sale in this state to an eligible product developer for purposes of
16 conducting testing required to support an application by such eligible
17 product developer for approval of a drug under subsections (b) and (j)
18 of Section 505 of the federal Food, Drug and Cosmetic Act, or the
19 licensing of a biological product under Section 351 of the federal Public
20 Health Service Act. Such manufacturer or wholesaler shall make the
21 drug available for sale to such eligible product developer at a price not
22 greater than the wholesale acquisition cost of the drug and without any
23 restriction that would block or delay the eligible product developer's
24 application in a manner inconsistent with Section 505-1(f)(8) of the
25 federal Food, Drug and Cosmetic Act.

26 (c) An eligible product developer that receives a drug at a price not
27 greater than the wholesale acquisition cost for such drug pursuant to
28 this section shall charge consumers in this state the same price or less
29 for the drug manufactured by such eligible product developer.

30 (d) A manufacturer or wholesaler registered under chapter 417 of the
31 general statutes shall not be liable for injuries alleged to have been
32 caused by the failure of the eligible product developer to include
33 adequate safety warnings on a product's label or by a defect in the
34 product's design if:

35 (1) Such manufacturer or wholesaler has made the product
36 distributed in this state available to an eligible product developer in
37 accordance with the provisions of this section; and

38 (2) The product was not manufactured or sold by such manufacturer
39 or wholesaler.

40 (e) A violation of any of the provisions of subsection (b) or (c) of this
41 section shall be deemed an unfair or deceptive trade practice under
42 subsection (a) of section 42-110b of the general statutes.

This act shall take effect as follows and shall amend the following sections:		
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Section 1	<i>October 1, 2022</i>	New section
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Statement of Purpose:

To promote competition in the prescription drug market by allowing developers of generic drugs and biosimilar products to obtain reference samples.

[Proposed deletions are enclosed in brackets. Proposed additions are indicated by underline, except that when the entire text of a bill or resolution or a section of a bill or resolution is new, it is not underlined.]