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2015-2016 Regular Sessions

IN ASSEMBLY

(PREFILED)

January 7, 2015

Introduced by M. of A. PAULIN, GOTTFRIED, DINOWITZ, GALEF, HOOPER, JAFFEE, MARKEY, MILLER, RIVERA -- read once and referred to the Committee on Health

AN ACT to amend the public health law and the education law, in relation to generic drug products; and to repeal paragraph (o) of subdivision 1 of section 206 of the public health law relating thereto

THE PEOPLE OF THE STATE OF NEW YORK, REPRESENTED IN SENATE AND ASSEMBLY, DO ENACT AS FOLLOWS:

- 1 Section 1. Paragraph (o) of subdivision 1 of section 206 of the public 2 health law is REPEALED.
 - S 2. The public health law is amended by adding a new section 280-a to read as follows:
 - S 280-A. GENERIC DRUG PRODUCTS. 1. THE COMMISSIONER SHALL ESTABLISH AND PUBLISH A LIST OF DRUG PRODUCTS, REFERRED TO IN THIS SECTION AS "GENERIC DRUG" PRODUCTS, EACH OF WHICH SHALL MEET THE FOLLOWING CONDITIONS:
 - (A) THE DRUG PRODUCT HAS BEEN CERTIFIED OR APPROVED BY THE COMMISSION-ER OF THE FEDERAL FOOD AND DRUG ADMINISTRATION AS BEING SAFE AND EFFEC-TIVE FOR ITS LABELED INDICATIONS FOR USE, AND A NEW-DRUG APPLICATION OR AN ABBREVIATED NEW-DRUG APPLICATION APPROVED PURSUANT TO THE FEDERAL FOOD, DRUG, AND COSMETIC ACT IS HELD FOR SUCH DRUG PRODUCT; AND
- (B) THE COMMISSIONER OF THE FEDERAL FOOD AND DRUG ADMINISTRATION 14 15 EVALUATED SUCH DRUG PRODUCT AS PHARMACEUTICALLY AND THERAPEUTICALLY EOUIVALENT AND HAS LISTED SUCH DRUG PRODUCT ON THE LIST OF APPROVED DRUG 16 PRODUCTS WITH THE THERAPEUTIC EQUIVALENCE EVALUATIONS, PROVIDED, 17 THAT THE LIST PREPARED BY THE COMMISSIONER SHALL NOT INCLUDE ANY 18 19 DRUG PRODUCT WHICH THE COMMISSIONER OF THE FEDERAL FOOD AND DRUG ADMIN-20 ISTRATION HAS IDENTIFIED AS HAVING AN ACTUAL OR POTENTIAL BIOEQUIVALENCE
- 21 PROBLEM.

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EXPLANATION--Matter in ITALICS (underscored) is new; matter in brackets [] is old law to be omitted.

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THE MANUFACTURER OF A GENERIC DRUG PRODUCT SHALL MAKE AVAILABLE TO BIOPHARMACEUTIC STUDIES THE DEPARTMENT THE AND SUMMARIES, INCIDENCE ADVERSE EVENTS, AND ASSOCIATED BIOEOUIVALENCE DATA AND OF ANALYTICAL METHODS, INCLUDING DISSOLUTION DATA AND TEST METHODS PROVIDED FEDERAL FOOD AND DRUG ADMINISTRATION AS PART OF THE APPLICATION FOR SUCH GENERIC DRUG PRODUCT. THE DEPARTMENT SHALL MAKE SUCH TION FREELY AND PUBLICLY AVAILABLE ON ITS WEBSITE.

- S 3. Paragraphs (a) and (d) of subdivision 6 of section 6810 of the education law, paragraph (a) as amended by chapter 590 of the laws of 2011 and paragraph (d) as added by chapter 913 of the laws of 1986, are amended to read as follows:
- 12 (a) Every prescription written in this state by a person authorized to 13 issue such prescription shall be on prescription forms containing one 14 line for the prescriber's signature. The prescriber's signature shall 15 validate the prescription. Every electronic prescription shall provide 16 for the prescriber's electronic signature, which shall validate the electronic prescription. Imprinted conspicuously on every prescription 17 18 written in this state in eight point upper case type immediately below 19 the signature line shall be the words: "THIS PRESCRIPTION WILL BE FILLED GENERICALLY UNLESS PRESCRIBER WRITES 'd a w' IN THE BOX BELOW". Unless 20 21 the prescriber writes daw in such box in the prescriber's own hand-22 writing or, in the case of electronic prescriptions, inserts an elec-23 tronic direction to dispense the drug as written, the prescriber's 24 signature or electronic signature shall designate approval of 25 tution by a pharmacist of a GENERIC drug product pursuant to [paragraph (o) of subdivision one of] section [two hundred six] TWO HUNDRED EIGHT-26 27 Y-A of the public health law. No other letters or marks in such box 28 shall prohibit substitution. No prescription forms used or intended to 29 be used by a person authorized to issue a prescription shall have 'd a 30 w' preprinted in such box. Such box shall be placed directly under the signature line and shall be three-quarters inch in length and one-half 31 32 inch in height, or in comparable form for an electronic prescription as 33 may be specified by regulation of the commissioner. Immediately below such box shall be imprinted in six point type the words 34 "Dispense As 35 Notwithstanding any other provision of law, no state official, agency, board or other entity shall promulgate any regulation or 36 guideline modifying those elements of the prescription form's contents 37 38 specified in this subdivision. To the extent otherwise permitted by law, 39 a prescriber may modify only those elements of the prescription form's 40 contents not specified in this subdivision. Notwithstanding any other provision of this section or any other law, when a generic drug is not 41 available and the brand name drug originally prescribed is available and 42 43 pharmacist agrees to dispense the brand name product for a price 44 that will not exceed the price that would have been charged for the 45 generic substitute had it been available, substitution of a generic drug product will not be required. If the generic drug product is not avail-46 47 able and a medical emergency situation, which for purposes 48 section is defined as any condition requiring alleviation of severe pain which threatens to cause disability or take life if not promptly 49 50 treated, exists, then the pharmacist may dispense the brand name product 51 at his regular price. In such instances the pharmacist must record the 52 date, hour and nature of the medical emergency on the back of the prescription and keep a copy of all such prescriptions. 53
 - (d) No prescriber shall be subjected to civil liability arising solely from authorizing, in accordance with this subdivision, the substitution by a pharmacist of a GENERIC drug product pursuant to [paragraph (o) of

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1 subdivision one of] section [two hundred six] TWO HUNDRED EIGHTY-A of
2 the public health law.
3 S 4. Paragraph (b) of subdivision 1 of section 6816-a of the education

- S 4. Paragraph (b) of subdivision 1 of section 6816-a of the education law, as added by chapter 776 of the laws of 1977, is amended to read as follows:
- (b) The substituted drug product is contained in the list of GENERIC drug products established pursuant to [paragraph (o) of subdivision one of] section [two hundred six] TWO HUNDRED EIGHTY-A of the public health law; and
- 10 S 5. This act shall take effect on the ninetieth day after it shall 11 have become a law. Effective immediately, the addition, amendment and/or 12 repeal of any rule or regulation necessary for the implementation of 13 this act on its effective date is authorized to be made and completed on 14 or before such effective date.