

7678--A

I N   S E N A T E

May 28, 2014

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Introduced by Sens. LIBOUS, VALESKY -- read twice and ordered printed,  
and when printed to be committed to the Committee on Higher Education  
-- committee discharged, bill amended, ordered reprinted as amended  
and recommitted to said committee

AN ACT to amend the education law, in relation to the practice of opto-  
metry

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

1     Section 1. Subdivision 1 of section 7101-a of the education law is  
2     amended by adding a new paragraph (g) to read as follows:  
3     (G) PHASE THREE THERAPEUTIC PHARMACEUTICAL AGENTS. PHASE THREE THERA-  
4     PEUTIC PHARMACEUTICAL AGENTS SHALL MEAN THOSE ORALLY ADMINISTERED DRUGS  
5     USED FOR THERAPEUTIC PURPOSES FOR THE TREATMENT OF DISEASES OF THE EYE  
6     AND ADNEXA AND SHALL BE LIMITED TO:  
7     (I) ANTIBIOTICS;  
8     (II) DECONGESTANTS/ANTI-ALLERGENIC/ANTI-HISTAMINES;  
9     (III) ANTIGLAUCOMAS; PROVIDED HOWEVER, WHEN PRESCRIBED OR ADMINISTERED  
10    FOR THE TREATMENT OF ACUTE ANGLE CLOSURE GLAUCOMA, THE PRESCRIBING OPTO-  
11    METRIST SHALL MAKE ALL REASONABLE EFFORTS IMMEDIATELY THEREAFTER TO  
12    REFER THE PATIENT TO A LICENSED PHYSICIAN SPECIALIZING IN DISEASES OF  
13    THE EYE AND PROVIDE NOTIFICATION IN ACCORDANCE WITH SUBDIVISION SIX-A OF  
14    THIS SECTION;  
15    (IV) ANTIVIRALS;  
16    (V) ONE THREE-DAY SUPPLY OF ANALGESICS, BUT SHALL NOT INCLUDE THOSE  
17    LISTED IN SCHEDULES I AND II OF THE UNIFORM CONTROLLED SUBSTANCES ACT;  
18    (VI) NONSTEROIDAL ANTI-INFLAMMATORY DRUGS;  
19    (VII) ONE FOURTEEN-DAY SUPPLY OF CORTICOSTEROIDS.  
20    S 2. Paragraphs (c) and (d) of subdivision 4 of section 7101-a of the  
21    education law are relettered paragraphs (d) and (e) and a new paragraph  
22    (c) is added to read as follows:  
23    (C) BEFORE USING OR PRESCRIBING PHASE THREE THERAPEUTIC PHARMACEUTICAL  
24    AGENTS, AN OPTOMETRIST MUST BE CERTIFIED TO PRESCRIBE DIAGNOSTIC PHARMA-  
25    CEUTICAL AGENTS AND PHASE ONE AND PHASE TWO THERAPEUTIC PHARMACEUTICAL  
26    AGENTS AND HAVE COMPLETED A THIRTY HOUR PHASE THREE THERAPEUTIC PHARMA-

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

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1 CEUTICAL AGENT CERTIFICATION COURSE, WITH A CURRICULUM DEVELOPED BY AN  
2 ACCREDITED COLLEGE OF OPTOMETRY IN COLLABORATION WITH A NEW YORK STATE  
3 ACCREDITED MEDICAL SCHOOL. THE CURRICULUM, WHICH SHALL BE APPROVED BY  
4 THE DEPARTMENT, SHALL INCLUDE, BUT NOT BE LIMITED TO, INSTRUCTION IN  
5 PHARMACOLOGY AND DRUG INTERACTION AND BE TAUGHT THROUGH CLINICAL CASE  
6 SCENARIOS AND EMPHASIZE CLINICAL DECISION MAKING. SUCH COURSE SHALL  
7 QUALIFY TOWARDS MEETING THE THIRTY-SIX HOURS OF CONTINUING EDUCATION PER  
8 TRIENNIAL REGISTRATION PERIOD REQUIRED BY SUBDIVISION SEVEN OF THIS  
9 SECTION. THIS REQUIREMENT FOR THE THIRTY HOUR PHASE THREE THERAPEUTIC  
10 PHARMACEUTICAL AGENT CERTIFICATION COURSE SHALL NOT APPLY TO THOSE OPTO-  
11 METRISTS WHO (I) GRADUATED FROM AN ACCREDITED COLLEGE OF OPTOMETRY  
12 SUBSEQUENT TO JANUARY FIRST, TWO THOUSAND FOUR AND (II) HAVE TAKEN AND  
13 SUCCESSFULLY PASSED EITHER THE TREATMENT AND MANAGEMENT OF OCULAR  
14 DISEASES PORTION OF THE NATIONAL BOARD OF EXAMINERS IN OPTOMETRY OR AN  
15 EXAMINATION ACCEPTABLE TO THE BOARD.

16 S 3. Subdivision 5 of section 7101-a of the education law, as added by  
17 chapter 517 of the laws of 1995, is amended to read as follows:

18 5. Suspension of certification. The department shall suspend the  
19 certification for the use and prescribing of phase one therapeutic  
20 agents of any optometrist who fails to receive certification for phase  
21 two therapeutic pharmaceutical agents within three years of having been  
22 certified for phase one therapeutic pharmaceutical agents. THE DEPART-  
23 MENT SHALL SUSPEND THE CERTIFICATION FOR THE USE AND PRESCRIBING OF  
24 PHASE ONE AND PHASE TWO THERAPEUTIC AGENTS OF ANY OPTOMETRIST WHO FAILS  
25 TO RECEIVE CERTIFICATION FOR PHASE THREE THERAPEUTIC PHARMACEUTICAL  
26 AGENTS WITHIN THREE YEARS OF APPROVAL BY THE DEPARTMENT OF A THIRTY HOUR  
27 PHASE THREE THERAPEUTIC PHARMACEUTICAL AGENT CERTIFICATION COURSE PURSU-  
28 ANT TO PARAGRAPH (C) OF SUBDIVISION FOUR OF THIS SECTION.

29 S 4. The opening paragraph of subdivision 6 of section 7101-a of the  
30 education law, as added by chapter 517 of the laws of 1995, is amended  
31 and two new subdivisions 6-a and 6-b are added to read as follows:

32 Consultation WITH USE OF CERTAIN PHASE TWO THERAPEUTIC PHARMACEUTICAL  
33 AGENTS.

34 6-A. NOTIFICATION OF USE OF PHASE THREE THERAPEUTIC PHARMACEUTICAL  
35 AGENTS. AN OPTOMETRIST SHALL, AS SOON AS PRACTICABLE, DOCUMENT IN THE  
36 PATIENT'S RECORD THE PRESCRIPTION OR USE OF PHASE THREE THERAPEUTIC  
37 PHARMACEUTICAL AGENTS. WITHIN SEVENTY-TWO HOURS OF PRESCRIBING OR USING  
38 A PHASE THREE THERAPEUTIC PHARMACEUTICAL AGENT, AN OPTOMETRIST SHALL  
39 NOTIFY THE PATIENT'S PRIMARY CARE PRACTITIONER, AND DOCUMENT SUCH  
40 NOTIFICATION WITH THE FOLLOWING INFORMATION:

41 (I) THE NAME OF SUCH AGENT;

42 (II) THE DOSE;

43 (III) THE FREQUENCY OF USE; AND

44 (IV) THE DURATION OF USE OR PRESCRIPTION.

45 6-B. CONSULTATION WITH USE OF PHASE THREE THERAPEUTIC PHARMACEUTICAL  
46 AGENTS. IF IN THE PROFESSIONAL JUDGMENT OF THE OPTOMETRIST, A PATIENT'S  
47 CONDITION DOES NOT RESULT IN AN ADEQUATE CLINICAL RESPONSE TO THE PHASE  
48 THREE THERAPEUTIC PHARMACEUTICAL AGENT THERAPY, THE OPTOMETRIST SHALL  
49 CONSULT WITH THE PATIENT'S PRIMARY CARE PRACTITIONER AS SOON AS CLIN-  
50 ICALLY PRUDENT.

51 S 5. Subdivision 7 of section 7101-a of the education law, as added by  
52 chapter 517 of the laws of 1995, is amended to read as follows:

53 7. Continuing education. Each optometrist certified to use phase one  
54 [or], phase two, OR PHASE THREE therapeutic pharmaceutical agents shall  
55 complete a minimum of thirty-six hours of continuing education per  
56 triennial registration period. The education shall be in the area of

1 ocular disease and pharmacology, AT LEAST SIX HOURS OF WHICH SHALL  
2 RELATE SPECIFICALLY TO SYSTEMIC DRUG USE AND INTERACTION, and may  
3 include both didactic and clinical components. Such educational programs  
4 shall be approved in advance by the department and evidence of the  
5 completion of this requirement shall be submitted with each application  
6 for license renewal as required by section sixty-five hundred two of  
7 this chapter.

8 S 6. The opening paragraph of subdivision 8 of section 7101-a of the  
9 education law, as added by chapter 517 of the laws of 1995, is amended  
10 to read as follows:

11 Notice to patient WITH USE OF CERTAIN PHASE TWO THERAPEUTIC PHARMACEU-  
12 TICAL AGENTS.

13 S 7. Subdivision 10 of section 7101-a of the education law, as added  
14 by chapter 517 of the laws of 1995, is amended to read as follows:

15 10. Pharmaceutical agents. Optometrists who have been approved and  
16 certified by the department shall be permitted to use the following  
17 drugs:

18 (a) Diagnostic pharmaceuticals.

19 (b) Those optometrists having been certified for phase one therapeutic  
20 pharmaceutical agents shall be authorized [(i) to use and recommend all  
21 nonprescription medications appropriate for ocular disease whether  
22 intended for topical or oral use; and (ii)] to use and prescribe all  
23 phase one therapeutic pharmaceutical agents which are FDA approved and  
24 commercially available.

25 In the event an optometrist treats a patient with topical antiviral or  
26 steroidal drugs and the patient's condition either fails to improve or  
27 worsens within five days, the optometrist shall notify a physician  
28 designated by the patient or, if none, by the treating optometrist.

29 (c) Those optometrists having been certified for phase two therapeutic  
30 pharmaceutical agents shall be authorized to use and prescribe phase two  
31 therapeutic pharmaceutical agents which are FDA approved and commercial-  
32 ly available.

33 (D) THOSE OPTOMETRISTS HAVING BEEN CERTIFIED FOR PHASE THREE THERAPEU-  
34 TIC PHARMACEUTICAL AGENTS SHALL BE AUTHORIZED TO USE AND PRESCRIBE PHASE  
35 THREE THERAPEUTIC PHARMACEUTICAL AGENTS WHICH ARE FDA APPROVED AND  
36 COMMERCIALY AVAILABLE.

37 (E) THOSE OPTOMETRISTS HAVING BEEN CERTIFIED FOR PHASE ONE, PHASE TWO  
38 OR PHASE THREE THERAPEUTIC PHARMACEUTICAL AGENTS SHALL BE AUTHORIZED TO  
39 USE AND RECOMMEND ALL NONPRESCRIPTION MEDICATIONS, WHETHER INTENDED FOR  
40 TOPICAL OR ORAL USE, APPROPRIATE FOR THE TREATMENT OF THE EYE AND  
41 ADNEXA.

42 S 8. Subdivision 12 of section 7101-a of the education law, as added  
43 by chapter 517 of the laws of 1995, is amended to read as follows:

44 12. Responsibilities of the commissioner of health. [The] IN ORDER TO  
45 SATISFY THE REQUIREMENT THAT AN OPTOMETRIST AUTHORIZED TO USE PHARMACEU-  
46 TICAL AGENTS FOR USE IN THE DIAGNOSIS, TREATMENT OR PREVENTION OF OCULAR  
47 DISEASE SHALL BE HELD TO THE SAME STANDARD OF CARE IN DIAGNOSIS, USE OF  
48 SUCH AGENTS, AND TREATMENT AS THAT DEGREE OF SKILL AND PROFICIENCY  
49 COMMONLY EXERCISED BY A PHYSICIAN IN THE SAME COMMUNITY, THE commission-  
50 er of health [may recommend to the commissioner additions or deletions  
51 to the department's regulations relating to optometric use of drugs  
52 except that such recommendations shall be limited only to additions  
53 which have been determined to be equivalent to those drugs already  
54 authorized or deletions based upon a finding that the drugs are no long-  
55 er appropriate for their current use or for other similar reasons.]  
56 SHALL RECOMMEND TO THE COMMISSIONER ADDITIONS OR DELETIONS TO THE LIST

1 OF APPROVED CATEGORIES OF DIAGNOSTIC, THERAPEUTIC AND ORAL PHARMACEU-  
2 TICAL AGENTS WHICH ARE CONSISTENT WITH THE STANDARD OF CARE IN THE  
3 COMMUNITY FOR THE DIAGNOSIS AND TREATMENT OF DISEASES OF THE EYE AND  
4 ADNEXA.

5 S 9. This act shall take effect on the one hundred twentieth day after  
6 it shall have become a law; provided that any rule or regulation neces-  
7 sary for the timely implementation of this act on its effective date  
8 shall be promulgated on or before such effective date.