

1 **SENATE FLOOR VERSION**

2 April 13, 2022

3 **AS AMENDED**

4 ENGROSSED HOUSE

5 BILL NO. 2649

6 By: Echols of the House

7 and

8 Garvin and Dugger of the
9 Senate

10
11 **[Oklahoma Durable Medical Equipment Licensing Act -
12 certain inspections - licensing qualifications -
13 codification]**

14 BE IT ENACTED BY THE PEOPLE OF THE STATE OF OKLAHOMA:

15 SECTION 1. NEW LAW A new section of law to be codified
16 in the Oklahoma Statutes as Section 375 of Title 59, unless there is
17 created a duplication in numbering, reads as follows:

18 This act shall be known and may be cited as the "Oklahoma
19 Durable Medical Equipment Licensing Act".

20 SECTION 2. NEW LAW A new section of law to be codified
21 in the Oklahoma Statutes as Section 376 of Title 59, unless there is
22 created a duplication in numbering, reads as follows:

23 As used in the Oklahoma Durable Medical Equipment Licensing Act:

24 1. "Board" means the State Board of Pharmacy;

2. a. "Durable medical equipment" means equipment for which
a prescription is required, including for repair and
replacement parts, and that:

- 1 (1) can stand repeated use,
- 2 (2) has an expected useful life of at least three (3)
- 3 years,
- 4 (3) is primarily and customarily used to serve a
- 5 medical purpose,
- 6 (4) is not generally useful to a person in the
- 7 absence of illness or injury,
- 8 (5) is appropriate for use in the home, and
- 9 (6) is intended for use by the consumer.

10 b. Durable medical equipment includes, but is not limited
11 to:

- 12 (1) ambulating assistance equipment,
- 13 (2) mobility equipment,
- 14 (3) rehabilitation seating,
- 15 (4) oxygen care and oxygen delivery systems,
- 16 (5) respiratory equipment and respiratory disease
- 17 management devices,
- 18 (6) rehabilitation environmental control equipment,
- 19 (7) ventilators,
- 20 (8) apnea monitors,
- 21 (9) diagnostic equipment,
- 22 (10) feeding pumps,
- 23 (11) beds prescribed by physicians to alleviate
- 24 medical conditions,

1 (12) transcutaneous electrical nerve stimulators, and

2 (13) sequential compression devices; and

3 3. "Supplier" means any person or entity that provides durable
4 medical equipment services or products and that currently bills or
5 plans to bill a claim for reimbursement of services or products to a
6 third party.

7 SECTION 3. NEW LAW A new section of law to be codified
8 in the Oklahoma Statutes as Section 377 of Title 59, unless there is
9 created a duplication in numbering, reads as follows:

10 A. Any supplier of durable medical equipment to a consumer in
11 Oklahoma shall possess a durable medical equipment supplier license
12 issued by the Board pursuant to this act.

13 B. Licenses issued by the Board pursuant to this act shall be
14 effective for twelve (12) months from the date of issuance and shall
15 not be transferable or assignable.

16 C. The Board shall have the authority to initially and
17 periodically inspect the applicant's office or place of business.

18 D. The Board shall promulgate rules necessary to implement the
19 provisions of this act. Such rules shall prioritize patient safety
20 and quality of durable medical equipment. The Board may provide by
21 rule that any person or entity accredited by organizations
22 recognized by the Centers for Medicare and Medicaid Services is
23 deemed to meet all or some of the requirements of this act.

1 E. Nothing in this section shall be construed to restrict or
2 prohibit private transactions between two parties.

3 SECTION 4. NEW LAW A new section of law to be codified
4 in the Oklahoma Statutes as Section 378 of Title 59, unless there is
5 created a duplication in numbering, reads as follows:

6 A. The Board shall be authorized to issue a license to an
7 applicant for licensure as a supplier of durable medical equipment
8 if the applicant:

9 1. Submits an application in a form prescribed by the Board;

10 2. Maintains a physical office or place of business within this
11 state;

12 3. Pays a license fee established by the Board;

13 4. Meets all state and federal accreditation requirements; and

14 5. Meets all safety standards established by the Board, which
15 shall include, but not be limited to:

16 a. ensuring that all personnel engaged in delivery,
17 maintenance and repair of durable medical equipment
18 receive annual continuing education,

19 b. instructing the patient or patient's caregiver about
20 how to use the durable medical equipment provided,

21 c. receiving and responding to complaints from patients,

22 d. maintaining records of all patients receiving durable
23 medical equipment, and

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1 e. managing, maintaining and servicing durable medical
2 equipment.

3 B. The Board may issue a license to a Medicare or Medicaid
4 enrolled out-of-state supplier who has at least one accredited
5 facility within one hundred (100) miles of any Oklahoma resident
6 being served by the supplier.

7 C. The Board may revoke or suspend a license for:

8 1. Violation of state or federal law;

9 2. Violation of rules promulgated pursuant to this act;

10 3. Permitting, aiding or abetting any illegal act;

11 4. Failing to meet the safety standards established by the
12 Board pursuant to this act;

13 5. Engaging in conduct or practices found by the Board to be
14 detrimental to the health, safety or welfare of patients; or

15 6. Failing to renew a license.

16 SECTION 5. NEW LAW A new section of law to be codified
17 in the Oklahoma Statutes as Section 379 of Title 59, unless there is
18 created a duplication in numbering, reads as follows:

19 The Oklahoma Durable Medical Equipment Licensing Act shall not
20 apply to:

21 1. Pharmacies and pharmacists;

22 2. Hospitals;

23 3. Ambulatory surgical centers;

- 1 4. Health care facilities owned or operated by the state or
2 federal government;
- 3 5. Skilled nursing facilities;
- 4 6. Assisted living facilities;
- 5 7. Prosthetic or orthotic practitioners;
- 6 8. Health care practitioners who are licensed to practice
7 health care in the State of Oklahoma and who provide durable medical
8 equipment within the scope of their health care practice;
- 9 9. Manufacturers or wholesale distributors that do not sell or
10 rent durable medical equipment directly to consumers;
- 11 10. Suppliers of insulin infusion pumps and related supplies or
12 services; **or**
- 13 **11. Suppliers of medical devices approved by the Food and Drug**
14 **Administration that are used in the treatment of cancerous tumors.**

15 COMMITTEE REPORT BY: COMMITTEE ON APPROPRIATIONS
16 April 13, 2022 - DO PASS AS AMENDED

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