

AMENDED IN ASSEMBLY APRIL 7, 2008
AMENDED IN ASSEMBLY MARCH 25, 2008
CALIFORNIA LEGISLATURE—2007–08 REGULAR SESSION

ASSEMBLY BILL

No. 2747

Introduced by Assembly Members Berg and Levine

February 22, 2008

An act to add Part 1.8 (commencing with Section 442) to Division 1 of the Health and Safety Code, relating to end-of-life care.

LEGISLATIVE COUNSEL'S DIGEST

AB 2747, as amended, Berg. End-of-life care.

Existing law provides for the licensure and regulation of health facilities and hospices by the State Department of Public Health. Existing law provides for the regulation and licensing of physicians and surgeons by the Medical Board of California.

This bill would provide that when an attending physician makes a diagnosis that a patient has a terminal illness or makes a prognosis that a patient has less than one year to live, the health care provider shall provide the patient with the opportunity to receive information and counseling regarding legal end-of-life options, as specified, and provide for the referral or transfer of a patient if the patient's physician does not wish to comply with the patient's choice of end-of-life options.

Vote: majority. Appropriation: no. Fiscal committee: no.
State-mandated local program: no.

The people of the State of California do enact as follows:

SECTION 1. The Legislature finds and declares all of the following:

(a) Palliative and hospice care are invaluable resources for terminally ill Californians in need of comfort and support at the end of life.

(b) Palliative care and conventional medical treatment should be thoroughly integrated rather than viewed as separate entities.

(c) Even though Californians with a prognosis of six months or less to live are eligible for hospice care, nearly two-thirds of them receive hospice services for less than one month.

(d) Many patients benefit from being referred to hospice care earlier, where they receive better pain and symptom management and have an improved quality of life.

(e) Significant information gaps may exist between health care providers and their patients on end-of-life care options potentially leading to delays to, or lack of, referrals to hospice care for terminally ill patients. The sharing of important information regarding specific treatment options in a timely manner by health care providers is a key component of quality end-of-life care. Information that is helpful to patients and their families includes, but is not limited to, the availability of hospice care, the efficacy and potential side effects of continued curative treatment, and withholding or withdrawal of life sustaining treatments.

(f) Terminally ill and dying patients rely on their health care providers to give them timely and informative data. Research shows a lack of communication between health care providers and their terminally ill patients can cause problems, including poor availability of, and lack of clarity regarding, advanced health care directives and patients' end-of-life care preferences. This lack of information and poor adherence to patient choices ~~results~~ result in "bad deaths" that cause needless physical and psychological suffering to patients and their families.

(g) Those problems are complicated by social issues, such as cultural and religious pressures for the providers, patients, and their family members. A recent survey found that providers that object to certain practices are less likely than others to believe they have an obligation to present all of the options to patients and refer patients to other providers, if necessary.

1 (h) Every medical school in California is required to include
2 end-of-life care issues in its curriculum and every physician in
3 California is required to complete continuing education courses
4 in end-of-life care.

5 (i) Palliative care is not a one-size-fits-all approach. Patients
6 have a range of diseases and respond differently to treatment
7 options. A key benefit of palliative care is that it customizes
8 treatment to meet the needs of each individual person.

9 (j) Informed patient choices will help terminally ill patients and
10 their families cope with one of life's most challenging situations.

11 SEC. 2. Part 1.8 (commencing with Section 442) is added to
12 Division 1 of the Health and Safety Code, to read:

13
14 PART 1.8. END-OF-LIFE CARE

15
16 442. For the purposes of this part, the following definitions
17 shall apply:

18 (a) "Curative treatment" means treatment intended to cure or
19 alleviate symptoms of a given disease or condition.

20 (b) "Hospice" means a specialized form of interdisciplinary
21 health care that is designed to provide palliative care, alleviate the
22 physical, emotional, social, and spiritual discomforts of an
23 individual who is experiencing the last phases of life due to the
24 existence of a terminal disease, and provide supportive care to the
25 primary caregiver and the family of the hospice patient, and that
26 meets all of the criteria specified in subdivision (b) of Section
27 1746.

28 (c) "Palliative care" means medical treatment, interdisciplinary
29 care, or consultation provided to a patient or family members, or
30 both, that has as its primary purpose the prevention of, or relief
31 from, suffering and the enhancement of the quality of life, rather
32 than treatment aimed at investigation and intervention for the
33 purpose of cure or prolongation of life as described in subdivision
34 (b) of Section 1339.31.

35 (d) "Palliative sedation" means the use of sedative medications
36 to relieve extreme suffering by making the patient unaware and
37 unconscious, while artificial food and hydration are withheld,
38 during the progression of the disease leading to the death of the
39 patient.

(e) “Refusal or withdrawal of life sustaining treatment” means forgoing treatment or medical procedures that replace or support an essential bodily function, including, but not limited to, cardiopulmonary resuscitation, mechanical ventilation, artificial nutrition and hydration, dialysis, and any other treatment or discontinuing any or all of those treatments after they have been used for a reasonable time.

(f) “Voluntary stopping of eating and drinking” or “VSED” means the voluntary refusal of a patient to eat and drink in order to alleviate his or her suffering, and includes the withholding or withdrawal of life-sustaining treatment at the request of the patient.

442.5. When an attending physician makes a diagnosis that a patient has a terminal illness or makes a prognosis that a patient has less than one year to live, the physician, ~~or in the case of a patient in a health facility, as defined in Section 1250, the health facility,~~ shall provide the patient with the opportunity to receive comprehensive information and counseling regarding legal end-of-life care options. *When a patient is in a health facility, as defined in Section 1250, the attending physician or medical director may refer the patient to a hospice provider or private or public agencies and community-based organizations that specialize in end-of-life care case management and consultation to receive information and counseling regarding legal end-of-life care options.*

(a) If the patient indicates a desire to receive the information and counseling, the information shall include, but not be limited to, the following:

(1) Hospice care at home or in a health care setting.

(2) A prognosis with and without the continuation of curative treatment.

(3) The patient’s right to refusal *of* or withdrawal from life-sustaining treatment.

(4) The patient’s right to continue to pursue curative treatment while receiving palliative care.

(5) The patient’s right to comprehensive pain and symptom management at the end of life, including, but not limited to, adequate pain medication, treatment of nausea, palliative chemotherapy, relief of shortness of breath and fatigue, VSED, and palliative sedation.

1 (b) The information described in subdivision (a) may, but is not
2 required to be, in writing.

3 (c) Counseling may include, but not be limited to, discussions
4 about the outcomes on the patient and his or her family, based on
5 the interest of the patient.

6 442.7. If a physician does not wish to comply with his or her
7 patient's choice of end-of-life options, the health care provider
8 shall do both of the following:

9 (a) Refer or transfer a patient to an alternative health care
10 provider.

11 (b) Provide the patient with information on procedures to
12 transfer to an alternative health care provider.