HOUSE No. 1526

The Commonwealth of Massachusetts

PRESENTED BY:

Jeffrey Sánchez

To the Honorable Senate and House of Representatives of the Commonwealth of Massachusetts in General Court assembled:

The undersigned legislators and/or citizens respectfully petition for the adoption of the accompanying bill:

An Act relative to the hemophilia advisory committee.

PETITION OF:

NAME:	DISTRICT/ADDRESS:	DATE ADDED:
Jeffrey Sánchez	15th Suffolk	2/4/2011
Viriato Manuel deMacedo	1st Plymouth	1/27/2011
Gale D. Candaras		2/4/2011
William N. Brownsberger		1/31/2011
Anne M. Gobi	5th Worcester	2/3/2011
Colleen M. Garry	36th Middlesex	2/3/2011
Jay R. Kaufman	15th Middlesex	2/3/2011
James B. Eldridge		2/3/2011
James Arciero	2nd Middlesex	2/4/2011

HOUSE No. 1526

By Mr. Sánchez of Boston, a petition (accompanied by bill, House, No. 1526) of Jeffrey Sánchez and others for legislation establishing a hemophilia advisory committee. Public Health.

The Commonwealth of Alassachusetts

In the Year Two Thousand Eleven

An Act relative to the hemophilia advisory committee.

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Be it enacted by the Senate and House of Representatives in General Court assembled, and by the authority of the same, as follows:

SECTION 1. Chapter 111 of the General Laws is hereby amended by striking out

sections 6B and 6C, as appearing in the 2008 Official Edition, and inserting in place thereof the

following sections:--

Section 6B. (a) There is hereby established a hemophilia advisory committee which shall review administrative actions and the impact of legislation relative to the treatment and care of persons with hemophilia or other inherited bleeding and clotting disorders and to make recommendations to the commissioner of public health and the commissioner of insurance relative to the establishment of programs for public awareness, reporting and treatment of hemophilia and other inherited bleeding and clotting disorders and to make recommendations for legislation relative thereto. The committee shall consist of 11 members. The commissioner of public health or a designee thereof and the commissioner of insurance or a designee thereof shall serve as non-voting members and co-chairs of the committee. The remaining members shall be appointed by the governor and include: 1 registered nurse, licensed, practicing and treating

individuals with hemophilia or other inherited bleeding and clotting disorders; 1 social worker, licensed, practicing and treating individuals with hemophilia or other inherited bleeding or clotting disorders; 1 representative of a commonwealth-based, federally funded hemophilia treatment center; 1 representative of a commonwealth provider of health insurance, established by statute; 1 representative of a voluntary health organization that services persons with hemophilia or other bleeding and clotting disorders; 1 person with hemophilia or a caregiver thereof; 1 person with a bleeding disorder other than hemophilia or a caregiver thereof; 1 person with a clotting disorder or a caregiver thereof; and 1 board certified physician, licensed, practicing and treating persons with hemophilia or other inherited bleeding and clotting disorders. Members of the committee shall receive no compensation but may be reimbursed for actual expenses incurred in carrying out their official duties. Six members shall constitute a quorum and the affirmative vote of six members shall be necessary for any action taken by the committee.

- (b) The hemophilia advisory committee shall make recommendations to the commissioners of public health and insurance relative to:
- (1) legislative or administrative changes to policies and programs integral to the health and wellness of persons with hemophilia or other inherited bleeding and clotting disorders;
- (2) coordination with other state agencies and private organizations to advance the development of community-based initiatives to increase the awareness of care and treatment of persons living with hemophilia or other inherited bleeding and clotting disorders;
- (3) dissemination of information relative to initiatives of interest to persons with hemophilia or other inherited bleeding and clotting disorders;

(4) standards of care and treatment for persons with hemophilia or other inherited bleeding and clotting disorders, taking into consideration the federal and state standards of care guidelines developed by state and national organizations including but not limited to the Medical and Scientific Advisory Council of the National Hemophilia Foundation;

- (5) protection and promotion of open access to any and all treatments for hemophilia and other inherited bleeding and clotting disorders consistent with the standards of care guidelines developed by state and national organizations including but not limited to the Medical and Scientific Advisory Council of the National Hemophilia Foundation;
- (6) assistance to persons living with hemophilia or other inherited bleeding and clotting disorders to access appropriate health insurance coverage through private or state-sponsored health insurance providers; and
- (7) development of, and participation in, care and treatment programs for persons with hemophilia or other inherited bleeding and clotting disorders, including self-administration, home care, medical and dental procedures and techniques designed to provide maximum control over bleeding episodes.
- (c) The department of public health may coordinate with other state agencies and private organizations, through cooperative agreements with hemophilia treatment centers, medical facilities, schools and nonprofit organizations that serve persons living with hemophilia or other inherited bleeding and clotting disorders, in order to advance the development of community based initiatives to increase the awareness of care and treatment for persons living with hemophilia or other inherited bleeding and clotting disorders. The department of public health

may also extend assistance to such agencies and organizations to facilitate connecting persons with hemophilia or other bleeding and clotting disorders with such agencies and programs.

Section 6C. The commissioner of public health, in consultation with the commissioner of insurance, shall annually submit a report to the joint committee on health care finance and the joint committee on public health on the status of care and treatment of persons with hemophilia or other inherited bleeding and clotting disorders. The report shall include information relative to the method and manner by which the commonwealth has protected the health and wellness of persons with hemophilia or other inherited bleeding and clotting disorders. A copy of said report shall also be submitted to the secretary of the executive health & human services and the director of the MassHealth program.

SECTION 2. The department of public health shall adopt such rules and regulations as necessary to effectuate the purposes of act.

SECTION 3. The initial report to be filed pursuant to section 2 shall be filed not later than December 31, 2011; and by inserting before the enacting clause the following emergency preamble:-

"Whereas, The deferred operation of this act would tend to defeat its purpose, which is to immediately reorganize the hemophilia advisory committee and authorize such committee to provide recommendations to the commonwealth relative to health insurance policies and health insurance plans and programs that impact individuals with hemophilia or other inherited bleeding and clotting disorders, therefore it is hereby declared to be an emergency law, necessary for the immediate preservation of the public convenience.