

AMENDED IN ASSEMBLY AUGUST 19, 2013

CALIFORNIA LEGISLATURE—2013–14 REGULAR SESSION

Assembly Joint Resolution

No. 28

Introduced by Assembly Member Brown

(Coauthors: Assembly Members Bradford, Hall, Holden, Jones-Sawyer, Mitchell, ~~and Weber~~ Weber, Achadjian, Alejo, Allen, Ammiano, Atkins, Bloom, Bocanegra, Bonilla, Bonta, Buchanan, Ian Calderon, Campos, Chau, Chávez, Chesbro, Conway, Cooley, Dahle, Daly, Dickinson, Eggman, Fong, Fox, Frazier, Beth Gaines, Garcia, Gatto, Gomez, Gonzalez, Gordon, Gorell, Gray, Grove, Hagman, Harkey, Roger Hernández, Jones, Levine, Linder, Logue, Lowenthal, Maienschein, Mansoor, Medina, Morrell, Mullin, Muratsuchi, Nazarian, Nestande, Pan, John A. Pérez, Quirk, Quirk-Silva, Rendon, Salas, Stone, Ting, Wagner, Waldron, Wieckowski, Wilk, Williams, and Yamada)

(Coauthor: Senator Wright)

July 3, 2013

Assembly Joint Resolution No. 28—Relative to Sickle Cell Anemia Awareness Month.

LEGISLATIVE COUNSEL'S DIGEST

AJR 28, as amended, Brown. Sickle Cell Anemia Awareness Month.

This measure would recognize the month of September 2013, and each September thereafter, as Sickle Cell Anemia Awareness Month in California, and would urge the Congress of the United States to support the President's continuation of funding for sickle cell anemia centers and research and to make sickle cell anemia and other genetic hemoglobin disorders a public health priority.

Fiscal committee: no.

1 WHEREAS, Sickle cell anemia and sickle cell disease, used
2 interchangeably, refer to a group of inherited disorders that affect
3 the red blood cells; and

4 WHEREAS, Sickle cell anemia is a disease in which a person's
5 body produces abnormally shaped red blood cells that resemble a
6 crescent or sickle, and which do not last as long as normal round
7 red blood cells, which leads to anemia. The sickle cells also get
8 stuck in blood vessels and block blood flow, which can cause pain
9 and organ damage; and

10 WHEREAS, Sickle cell anemia is a genetic disorder where
11 individuals with the disease are born with two sickle cell genes,
12 each inherited from one parent. An individual with only one sickle
13 cell gene has a "sickle cell trait," which occurs in one out of every
14 12 African Americans and in one out of every 100 Latinos in the
15 United States; and

16 WHEREAS, Unlike most people with sickle cell anemia, most
17 people who have a sickle cell trait never know they have it and
18 can live their entire lives without any complications from it; and

19 WHEREAS, Serious problems associated with a sickle cell trait
20 are rare. However, exercise-related sudden death in individuals
21 who have a sickle cell trait most commonly occurs in those
22 undergoing intense physical exertion, such as military recruits in
23 basic training and athletes during conditioning workouts; and

24 WHEREAS, Individuals with a sickle cell trait should not be
25 excluded from physical activity, including sports, unless
26 recommended to by medical personnel. Instead, people should be
27 educated about precautions that should be taken, including drinking
28 adequate amounts of fluids, pacing training with longer periods
29 of rest and recovery, avoiding participation in performance tests
30 such as sprints and mile runs, and, most importantly, being familiar
31 with the symptoms of overexertion; and

32 WHEREAS, It is estimated that more than 90,000 Americans
33 have sickle cell anemia. Sickle cell anemia occurs in one out of
34 every 500 African American births and in one out of every 36,000
35 Latino births; and

36 WHEREAS, Sickle cell anemia can be a life-threatening
37 condition, and access to comprehensive care can be limited by
38 social, economic, cultural, and geographic barriers; and

39 WHEREAS, The average cost of hospitalization for sickle cell
40 anemia in 2004 was \$6,223, for more than 84,000 hospital

1 admissions that year. Total hospitalization costs for individuals
2 with sickle cell anemia equaled \$488,000,000, of which 65 percent
3 were covered by Medicaid funds; and

4 WHEREAS, Individuals living with sickle cell anemia encounter
5 barriers to obtaining quality care and improving their quality of
6 life. These barriers include limitations in geographic access to
7 comprehensive care, the varied use of effective treatments, the
8 high reliance on emergency departments and on public health
9 programs, and the limited number of health care providers with
10 knowledge and experience to manage and treat sickle cell anemia;
11 and

12 WHEREAS, The Sickle Cell Anemia Control Act was signed
13 into law in 1972 by President Richard Nixon after pledging that
14 his administration would “reverse the record of neglect on the
15 dreaded disease” by increasing funding for and expanding sickle
16 cell anemia-related programs, including the development of
17 comprehensive sickle cell anemia centers; and

18 WHEREAS, In 1975, the Sickle Cell Disease Association of
19 America, Inc. and its member organizations began conducting
20 monthlong events in September to call attention to sickle cell
21 anemia and the need to address the problem at national and local
22 levels, and chose September as National Sickle Cell Awareness
23 Month in order for the public to reflect on the children and adults
24 whose lives, education, and careers have been affected by this
25 disease; and

26 WHEREAS, In 2003, the Sickle Cell Treatment Act was signed
27 into law; and

28 WHEREAS, The effort to officially recognize Sickle Cell
29 Anemia Awareness Month succeeded at the federal level in 1983
30 when the United States House of Representatives unanimously
31 passed, and President Ronald Reagan signed, the first resolution
32 introduced by the Congressional Black Caucus that recognized
33 September as National Sickle Cell Anemia Awareness Month;
34 now, therefore, be it

35 *Resolved by the Assembly and the Senate of the State of*
36 *California, jointly,* That the Legislature recognizes September
37 2013, and each September thereafter, as Sickle Cell Anemia
38 Awareness Month; and be it further

39 *Resolved,* That the Legislature urges the Congress of the United
40 States to support the President’s continuation of funding for the

1 Sickle Cell Disease Treatment Demonstration Program, the
2 Registry and Surveillance System for Hemoglobinopathy Program
3 Initiative, and the Public Health Approach Disorders program, and
4 to make sickle cell anemia and other genetic hemoglobin disorders
5 a public health priority; and be it further
6 *Resolved*, That the Chief Clerk of the Assembly transmit copies
7 of this resolution to the President and Vice President of the United
8 States, the Speaker of the United States House of Representatives,
9 the President pro Tempore of the United States Senate, each
10 Senator and Representative from California in the Congress of the
11 United States, and to the author for appropriate distribution.

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