

Senators on both sides of the aisle to support this important legislation to help this country take another step toward reclaiming our place as a guardian of human rights.

SUBMITTED RESOLUTIONS

SENATE RESOLUTION 202—TO PROVIDE FOR ISSUANCE OF A SUMMONS AND FOR RELATED PROCEDURES CONCERNING THE ARTICLES OF IMPEACHMENT AGAINST SAMUEL B. KENT

Mr. REID (for himself and Mr. MCCONNELL) submitted the following resolution; which was considered and agreed to:

S. RES. 202

Resolved, That a summons shall be issued which commands Samuel B. Kent to file with the Secretary of the Senate an answer to the articles of impeachment no later than July 2, 2009, and thereafter to abide by, obey, and perform such orders, directions, and judgments as the Senate shall make in the premises, according to the Constitution and laws of the United States.

SEC. 2. The Sergeant at Arms is authorized to utilize the services of the Deputy Sergeant at Arms or another employee of the Senate in serving the summons.

SEC. 3. The Secretary shall notify the House of Representatives of the filing of the answer and shall provide a copy of the answer to the House.

SEC. 4. The Managers on the part of the House may file with the Secretary of the Senate a replication no later than July 7, 2009.

SEC. 5. The Secretary shall notify counsel for Samuel B. Kent of the filing of a replication, and shall provide counsel with a copy.

SEC. 6. The Secretary shall provide the answer and the replication, if any, to the Presiding Officer of the Senate on the first day the Senate is in session after the Secretary receives them, and the Presiding Officer shall cause the answer and replication, if any, to be printed in the Senate Journal and in the Congressional Record. If a timely answer has not been filed, the Presiding Officer shall cause a plea of not guilty to be entered.

SEC. 7. The articles of impeachment, the answer, and the replication, if any, together with the provisions of the Constitution on impeachment, and the Rules of Procedure and Practice in the Senate When Sitting on Impeachment Trials, shall be printed under the direction of the Secretary as a Senate document.

SEC. 8. The provisions of this resolution shall govern notwithstanding any provisions to the contrary in the Rules of Procedure and Practice in the Senate When Sitting on Impeachment Trials.

SEC. 9. The Secretary shall notify the House of Representatives of this resolution.

SENATE RESOLUTION 203—TO PROVIDE FOR THE APPOINTMENT OF A COMMITTEE TO RECEIVE AND TO REPORT EVIDENCE WITH RESPECT TO ARTICLES OF IMPEACHMENT AGAINST JUDGE SAMUEL B. KENT

Mr. REID (for himself and Mr. MCCONNELL) submitted the following resolution; which was considered and agreed to:

S. RES. 203

Resolved, That pursuant to Rule XI of the Rules of Procedure and Practice in the Senate When Sitting on Impeachment Trials, the Presiding Officer shall appoint a committee of twelve senators to perform the duties and to exercise the powers provided for in the rule.

SEC. 2. The majority and minority leader shall each recommend six members and a chairman and vice chairman respectively to the Presiding Officer for appointment to the committee.

SEC. 3. The committee shall be deemed to be a standing committee of the Senate for the purpose of reporting to the Senate resolutions for the criminal or civil enforcement of the committee's subpoenas or orders, and for the purpose of printing reports, hearings, and other documents for submission to the Senate under Rule XI.

SEC. 4. During proceedings conducted under Rule XI the chairman of the committee is authorized to waive the requirement under the Rules of Procedure and Practice in the Senate When Sitting on Impeachment Trials that questions by a Senator to a witness, a manager, or counsel shall be reduced to writing and put by the Presiding Officer.

SEC. 5. In addition to a certified copy of the transcript of the proceedings and testimony had and given before it, the committee is authorized to report to the Senate a statement of facts that are uncontested and a summary, with appropriate references to the record, of evidence that the parties have introduced on contested issues of fact.

SEC. 6. The actual and necessary expenses of the committee, including the employment of staff at an annual rate of pay, and the employment of consultants with prior approval of the Committee on Rules and Administration at a rate not to exceed the maximum daily rate for a standing committee of the Senate, shall be paid from the contingent fund of the Senate from the appropriation account "Miscellaneous Items" upon vouchers approved by the chairman of the committee, except that no voucher shall be required to pay the salary of any employee who is compensated at an annual rate of pay.

SEC. 7. The Committee appointed pursuant to section one of this resolution shall terminate no later than 45 days after the pronouncement of judgment by the Senate on the articles of impeachment.

SEC. 8. The Secretary shall notify the House of Representatives and counsel for Judge Samuel B. Kent of this resolution.

SENATE RESOLUTION 204—DESIGNATING MARCH 31, 2010, AS "NATIONAL CONGENITAL DIAPHRAGMATIC HERNIA AWARENESS DAY"

Mr. VITTER submitted the following resolution; which was referred to the Committee on the Judiciary:

S. RES. 204

Whereas the congenital diaphragmatic hernia birth defect is one of the most prevalent, life-threatening birth defects in the United States;

Whereas the congenital diaphragmatic hernia birth defect is a severe, often deadly birth defect that has a devastating impact, in both human and economic terms, affecting equally people of all races, sexes, nationalities, geographic locations, and income levels;

Whereas the congenital diaphragmatic hernia birth defect occurs in 1 in every 2,000 live births in the United States and accounts for 8 percent of all major congenital anomalies;

Whereas, in 2004, there were approximately 4,115,590 live births in the United States, and in approximately 1,800 of those live births, the congenital diaphragmatic hernia birth defect occurred, causing countless additional friends, loved ones, spouses, and caregivers to shoulder the physical, emotional, and financial burdens the congenital diaphragmatic hernia birth defect causes;

Whereas there is no genetic indicator or any other indicator available to predict the occurrence of the congenital diaphragmatic hernia birth defect, other than through the performance of an ultrasound during pregnancy;

Whereas there is no consistent treatment or cure for the congenital diaphragmatic hernia birth defect;

Whereas the congenital diaphragmatic hernia birth defect is a leading cause of neonatal death in the United States;

Whereas 50 percent of the patients who do survive the congenital diaphragmatic hernia birth defect have residual health issues, resulting in a severe strain on pediatric medical resources and on the delivery of health care services in the United States;

Whereas proactive diagnosis and the appropriate management and care of fetuses afflicted with the congenital diaphragmatic hernia birth defect minimize the incidence of emergency situations resulting from the birth defect and dramatically improve survival rates among people with the birth defect;

Whereas neonatal medical care is one of the most expensive types of medical care provided in the United States and patients with the congenital diaphragmatic hernia birth defect stay in intensive care for approximately 60 to 90 days, costing millions of dollars, utilizing blood from local blood banks, and requiring the most technically advanced medical care;

Whereas the congenital diaphragmatic hernia birth defect is a birth defect that causes damage to the lungs and the cardiovascular system;

Whereas patients with the congenital diaphragmatic hernia birth defect may have long-term health issues such as respiratory insufficiency, gastroesophageal reflux, poor growth, neurodevelopmental delay, behavior problems, hearing loss, hernia recurrence, and orthopedic deformities;

Whereas the severity of the symptoms and outcomes of the congenital diaphragmatic hernia birth defect and the limited public awareness of the birth defect cause many patients to receive substandard care, to forego regular visits to physicians, and not to receive good health or therapeutic management that would help avoid serious complications in the future, compromising the quality of life of those patients;

Whereas people suffering from chronic, life-threatening diseases and birth defects, similar to the congenital diaphragmatic hernia birth defect, and family members of those people are predisposed to depression and the resulting consequences of depression because of anxiety over the possible pain, suffering, and premature death that people with such diseases and birth defects may face;

Whereas the Senate and taxpayers of the United States want treatments and cures for disease and hope to see results from investments in research conducted by the National Institutes of Health and from initiatives such as the National Institutes of Health Roadmap to the Future;

Whereas the congenital diaphragmatic hernia birth defect is an example of how collaboration, technological innovation, scientific momentum, and public-private partnerships can generate therapeutic interventions that directly benefit the people and

families suffering from the congenital diaphragmatic hernia birth defect;

Whereas collaboration, technological innovation, scientific momentum, and public-private partnerships can save billions of Federal dollars under Medicare, Medicaid, and other programs for therapies, and early intervention will increase survival rates among people suffering from the congenital diaphragmatic hernia birth defect;

Whereas improvements in diagnostic technology, the expansion of scientific knowledge, and better management of care for patients with the congenital diaphragmatic hernia birth defect already have increased survival rates in some cases;

Whereas there is still a need for more research and increased awareness of the congenital diaphragmatic hernia birth defect and for an increase in funding for that research in order to provide a better quality of life to survivors of the congenital diaphragmatic hernia birth defect, and more optimism for the families and health care professionals who work with children with the birth defect;

Whereas there are thousands of volunteers nationwide dedicated to expanding research, fostering public awareness and understanding, educating patients and their families about the congenital diaphragmatic hernia birth defect to improve their treatment and care, providing appropriate moral support, and encouraging people to become organ donors; and

Whereas volunteers engage in an annual national awareness event held on March 31, making that day an appropriate time to recognize National Congenital Diaphragmatic Hernia Awareness Day: Now, therefore, be it

Resolved, That the Senate—

(1) designates March 31, 2010, as “National Congenital Diaphragmatic Hernia Awareness Day”;

(2) supports the goals and ideals of a national day to raise public awareness and understanding of the congenital diaphragmatic hernia birth defect;

(3) recognizes the need for additional research into a cure for the congenital diaphragmatic hernia birth defect; and

(4) encourages the people of the United States and interested groups to support National Congenital Diaphragmatic Hernia Awareness Day through appropriate ceremonies and activities, to promote public awareness of the congenital diaphragmatic hernia birth defect, and to foster understanding of the impact of the disease on patients and their families.

SENATE RESOLUTION 205—SUPPORTING THE GOALS AND IDEALS OF AFRICAN AMERICAN BONE MARROW AWARENESS MONTH

Ms. STABENOW (for herself and Mr. ISAKSON) submitted the following resolution; which was considered and agreed to:

S. RES. 205

Whereas a bone marrow or blood cell transplant is a potentially life-saving treatment for patients with leukemia, lymphoma, and other blood diseases;

Whereas a bone marrow or blood cell transplant replaces a patient's unhealthy blood cells with healthy blood-forming cells from a volunteer donor;

Whereas a patient who does not have a suitably matching donor in the family may search the National Marrow Donor Program Donor Registry for a donor;

Whereas blood or cell samples from adult donors or cord blood units are tested and the

tissue or cell type is added to the National Marrow Donor Program Donor Registry, and physicians may search that registry when they need to find donors whose tissue type matches their patients’;

Whereas African Americans make up 8 percent of, or more than 550,000 of the 7,000,000 people currently on, the National Marrow Donor Program Donor Registry;

Whereas of the 35,000 people that have received transplants since the inception of the National Marrow Donor Program Donor Registry, only 1,500 have been African Americans;

Whereas more than 70 life-threatening diseases can be treated with a bone marrow transplant;

Whereas there is a possibility that an African American patient could match a donor from any racial or ethnic group, but the most likely match is another African American;

Whereas to become a volunteer donor, potential donors must be between 18 and 60 years of age, meet health guidelines, provide a small blood sample or swab of cheek cells to determine the donor's tissue type, complete a brief health questionnaire, and sign a consent form to have the tissue type of the donor listed on the Donor Registry;

Whereas the Bone Marrow Wish Organization, which is a minority-run nonprofit organization based in Detroit that was started by an actual bone marrow donor, is initiating “African American Bone Marrow Awareness Month”;

Whereas the annual month of awareness would promote donor awareness and increase the number of African Americans registered with the National Marrow Donor Program throughout the Nation; and

Whereas July 2009 would be an appropriate month to observe African American Bone Marrow Awareness Month: Now, therefore, be it

Resolved, That the Senate—

(1) supports the goals and ideals of African American Bone Marrow Awareness Month;

(2) urges the people of the United States to participate in appropriate programs and activities with respect to bone marrow awareness, including speaking with health care professionals about bone marrow donation; and

(3) urges all people of the United States to register to become blood marrow donors and encourages all people of the United States to organize blood marrow registration drives in their communities.

NOTICE OF HEARING

IMPEACHMENT TRIAL COMMITTEE ON THE ARTICLES AGAINST JUDGE SAMUEL B. KENT

Ms. MCCASKILL. Mr. President, I wish to announce that the Impeachment Trial Committee on the Articles Against Judge Samuel B. Kent will meet on Thursday, June 25, 2009, at 4:30 p.m., to conduct its organization meeting.

For further information regarding this meeting, please contact Peg Gustafson on 202-224-6154.

AUTHORITY FOR COMMITTEES TO MEET

COMMITTEE ON ARMED SERVICES

Mr. CARDIN. Mr. President, I ask unanimous consent that the Committee on Armed Services be authorized to meet during the session of the Senate on Wednesday, June 24, 2009, at 2:30 p.m.

The PRESIDING OFFICER. Without objection, it is so ordered.

COMMITTEE ON COMMERCE, SCIENCE, AND TRANSPORTATION

Mr. CARDIN. Mr. President, I ask unanimous consent that the Committee on Commerce, Science, and Transportation be authorized to meet during the session of the Senate on Wednesday, June 24, 2009, at 2:30 p.m., in room 253 of the Russell Senate Office Building.

The PRESIDING OFFICER. Without objection, it is so ordered.

COMMITTEE ON ENVIRONMENT AND PUBLIC WORKS

Mr. CARDIN. Mr. President, I ask unanimous consent that the Committee on Environment and Public Works be authorized to meet during the session of the Senate on Wednesday, June 24, 2009 at 10:45 a.m. in room 406 of the Dirksen Senate Office Building.

The PRESIDING OFFICER. Without objection, it is so ordered.

COMMITTEE ON FOREIGN RELATIONS

Mr. CARDIN. Mr. President, I ask unanimous consent that the Committee on Foreign Relations be authorized to meet during the session of the Senate on Wednesday, June 24, 2009, at 11 a.m., to hold a roundtable entitled “Iran at a Crossroads?”.

The PRESIDING OFFICER. Without objection, it is so ordered.

COMMITTEE ON FOREIGN RELATIONS

Mr. CARDIN. Mr. President, I ask unanimous consent that the Committee on Foreign Relations be authorized to meet during the session of the Senate on Wednesday, June 24, 2009, at 2:30 p.m.

The PRESIDING OFFICER. Without objection, it is so ordered.

COMMITTEE ON HEALTH, EDUCATION, LABOR, AND PENSIONS

Mr. CARDIN. Mr. President, I ask unanimous consent that the Committee on Health, Education, Labor, and Pensions be authorized to meet during the session of the Senate on Wednesday, June 24, 2009, at 10 a.m. in room 325 of the Russell Senate Office Building.

The PRESIDING OFFICER. Without objection, it is so ordered.

COMMITTEE ON HOMELAND SECURITY AND GOVERNMENT AFFAIRS

Mr. CARDIN. Mr. President, I ask unanimous consent that the Committee on Homeland Security and Governmental Affairs be authorized to meet during the session of the Senate on Wednesday, June 24, 2009, at 9 a.m. to conduct a hearing entitled “Type 1 Diabetes Research: Real Progress and Real Hope for a Cure.”

The PRESIDING OFFICER. Without objection, it is so ordered.

COMMITTEE ON THE JUDICIARY

Mr. CARDIN. Mr. President, I ask unanimous consent that the Committee on the Judiciary be authorized to meet during the session of the Senate on Wednesday, June 24, 2009, at 10 a.m., in room SD-226 of the Dirksen