

S. 694, a bill to provide assistance to Best Buddies to support the expansion and development of mentoring programs, and for other purposes.

S. 936

At the request of Mr. LAUTENBERG, the name of the Senator from Rhode Island (Mr. REED) was added as a cosponsor of S. 936, a bill to amend the Federal Water Pollution Control Act to authorize appropriations for sewer overflow control grants.

S. 1058

At the request of Mr. UDALL of Colorado, the names of the Senator from Georgia (Mr. ISAKSON) and the Senator from Florida (Mr. LEMIEUX) were added as cosponsors of S. 1058, a bill to amend the Internal Revenue Code of 1986 to reduce the tax on beer to its pre-1991 level, and for other purposes.

S. 1111

At the request of Mr. ROCKEFELLER, the name of the Senator from Massachusetts (Mr. KERRY) was added as a cosponsor of S. 1111, a bill to require the Secretary of Health and Human Services to enter into agreements with States to resolve outstanding claims for reimbursement under the Medicare program relating to the Special Disability Workload project.

S. 1234

At the request of Mr. LIEBERMAN, the name of the Senator from Florida (Mr. LEMIEUX) was added as a cosponsor of S. 1234, a bill to modify the prohibition on recognition by United States courts of certain rights relating to certain marks, trade names, or commercial names.

S. 1329

At the request of Mr. KOHL, the name of the Senator from South Dakota (Mr. JOHNSON) was added as a cosponsor of S. 1329, a bill to authorize the Attorney General to award grants to State courts to develop and implement State courts interpreter programs.

S. 1345

At the request of Mr. REED, the name of the Senator from Alaska (Mr. BEGICH) was added as a cosponsor of S. 1345, a bill to aid and support pediatric involvement in reading and education.

S. 1859

At the request of Mr. ROCKEFELLER, the name of the Senator from Oregon (Mr. MERKLEY) was added as a cosponsor of S. 1859, a bill to reinstate Federal matching of State spending of child support incentive payments.

S. 2760

At the request of Mr. UDALL of New Mexico, the name of the Senator from New Mexico (Mr. BINGAMAN) was added as a cosponsor of S. 2760, a bill to amend title 38, United States Code, to provide for an increase in the annual amount authorized to be appropriated to the Secretary of Veterans Affairs to carry out comprehensive service programs for homeless veterans.

S. 2796

At the request of Mr. ENZI, the name of the Senator from Arkansas (Mrs.

LINCOLN) was added as a cosponsor of S. 2796, a bill to extend the authority of the Secretary of Education to purchase guaranteed student loans for an additional year, and for other purposes.

S. 2853

At the request of Mr. CRAPO, his name was withdrawn as a cosponsor of S. 2853, a bill to establish a Bipartisan Task Force for Responsible Fiscal Action, to assure the long-term fiscal stability and economic security of the Federal Government of the United States, and to expand future prosperity growth for all Americans.

At the request of Mr. BROWNBACK, his name was withdrawn as a cosponsor of S. 2853, *supra*.

S. 2885

At the request of Ms. LANDRIEU, the name of the Senator from Wisconsin (Mr. FEINGOLD) was added as a cosponsor of S. 2885, a bill to amend the Omnibus Crime Control and Safe Streets Act of 1968 to provide adequate benefits for public safety officers injured or killed in the line of duty, and for other purposes.

S. 2908

At the request of Mr. KOHL, the name of the Senator from Michigan (Ms. STABENOW) was added as a cosponsor of S. 2908, a bill to amend the Energy Policy and Conservation Act to require the Secretary of Energy to publish a final rule that establishes a uniform efficiency descriptor and accompanying test methods for covered water heaters, and for other purposes.

S. 2926

At the request of Mrs. LINCOLN, the name of the Senator from Pennsylvania (Mr. CASEY) was added as a cosponsor of S. 2926, a bill to amend the XVIII of the Social Security Act to provide for the application of a consistent Medicare part B premium for all Medicare beneficiaries in a budget neutral manner for 2010, to provide an additional round of economic recovery payments to certain beneficiaries, and to assess the need for a consumer price index for elderly consumers to compute cost-of-living increases for certain governmental benefits.

S. 2936

At the request of Mr. BAUCUS, the names of the Senator from California (Mrs. FEINSTEIN), the Senator from Maryland (Mr. CARDIN), the Senator from Alaska (Mr. BEGICH), the Senator from Montana (Mr. TESTER), the Senator from Missouri (Mr. BOND), the Senator from Kansas (Mr. ROBERTS), the Senator from Tennessee (Mr. ALEXANDER), the Senator from South Dakota (Mr. THUNE), the Senator from Colorado (Mr. BENNET) and the Senator from Vermont (Mr. LEAHY) were added as cosponsors of S. 2936, a bill to accelerate the income tax benefits for charitable cash contributions for the relief of victims of the earthquake in Haiti.

S. 2938

At the request of Mr. THUNE, the names of the Senator from Kansas (Mr. ROBERTS), the Senator from Maine (Ms.

COLLINS), the Senator from Utah (Mr. HATCH), the Senator from Idaho (Mr. RISCH), the Senator from Georgia (Mr. ISAKSON), the Senator from Arizona (Mr. KYL) and the Senator from Alaska (Ms. MURKOWSKI) were added as cosponsors of S. 2938, a bill to terminate authority under the Troubled Asset Relief Program, and for other purposes.

S. CON. RES. 39

At the request of Mr. MENENDEZ, the name of the Senator from Minnesota (Mr. FRANKEN) was added as a cosponsor of S. Con. Res. 39, a concurrent resolution expressing the sense of the Congress that stable and affordable housing is an essential component of an effective strategy for the prevention, treatment, and care of human immunodeficiency virus, and that the United States should make a commitment to providing adequate funding for the development of housing as a response to the acquired immunodeficiency syndrome pandemic.

S. RES. 373

At the request of Mr. CRAPO, the names of the Senator from Idaho (Mr. RISCH) and the Senator from Mississippi (Mr. COCHRAN) were added as cosponsors of S. Res. 373, a resolution designating the month of February 2010 as "National Teen Dating Violence Awareness and Prevention Month".

AMENDMENT NO. 3301

At the request of Mr. THUNE, the names of the Senator from Arizona (Mr. KYL), the Senator from Alaska (Ms. MURKOWSKI), the Senator from Utah (Mr. HATCH), the Senator from Wisconsin (Mr. FEINGOLD), the Senator from Georgia (Mr. CHAMBLISS), the Senator from Idaho (Mr. RISCH), the Senator from Maine (Ms. COLLINS), the Senator from Kansas (Mr. ROBERTS), the Senator from New Hampshire (Mr. GREGG), the Senator from Montana (Mr. TESTER), the Senator from Georgia (Mr. ISAKSON) and the Senator from Tennessee (Mr. CORKER) were added as cosponsors of amendment No. 3301 proposed to H.J. Res. 45.

STATEMENTS ON INTRODUCED BILLS AND JOINT RESOLUTIONS

By Mr. PRYOR (for himself and Mr. CARDIN):

S. 2942. A bill to amend the Federal Food, Drug, and Cosmetic Act to establish a nanotechnology program; to the Committee on Health, Education, Labor, and Pensions.

Mr. PRYOR. Mr. President, I rise today with Senator CARDIN to introduce the Nanotechnology Safety Act of 2010 which will authorize a program of scientific investigation by the Food and Drug Administration on nanotechnology-based medical and health products.

Nanotechnology holds great promise to revolutionize the development of new medicines, drug delivery, and orthopedic implants while holding down the cost of health care. However, Congress and the FDA must assure the

public that nanotechnology-based products are both safe and efficacious. The Nanotechnology Safety Act of 2010 will enable the FDA to properly study how nanomaterials are absorbed by the human body, how nanomaterials designed to carry cancer fighting drugs target and kill tumors, and how nanoscale texturing of bone implants can make a stronger joint and reduce the threat of infection.

Nanotechnology, or the manipulation of material at dimensions between 1 and 100 nanometers, is a challenging scientific area. To put this size scale in perspective, a human hair is 80,000 nanometers thick.

Nanomaterials have different chemical, physical, electrical and biological characteristics than when used as larger, bulk materials. For example, nanoscale silver has exhibited unique antibacterial properties for treating infections and wounds. Nanomaterials have a much larger ratio of surface area to mass than ordinary materials do. It is at the surface of materials that biological and chemical reactions take place, and so we would expect nanomaterials to be more reactive than bulk materials.

The novel characteristics of nanomaterials mean that risk assessments developed for ordinary materials may be of limited use in determining the health and public safety of products based on nanotechnology.

The FDA needs the tools and resources to assure the public that nanotechnology-based medical and health products are safe and effective. The development of a regulatory framework for the use of nanomaterials in drugs, medical devices, and food additives must be based on scientific knowledge and data about each specific technology and product. Without a robust scientific framework there is no way to know what data to collect. More than a dozen material characteristics have been suggested even for relatively simple nanomaterials. Without better scientific knowledge of nanomaterials and their behavior in the human body, we do not know what data to collect and examine.

In 2007, the FDA Nanotechnology Task Force published a report analyzing the FDA's scientific program and regulatory authority for addressing nanotechnology in drugs, medical devices, biologics, and food supplements. A general finding of the report is that nanoscale materials present regulatory challenges similar to those posed by products using other emerging technologies. However, these challenges may be magnified because nanotechnology can be used to make almost any FDA-regulated product. Also, at the nanoscale, the properties of a material relevant to the safety and effectiveness of the FDA-regulated products might change.

The Task Force recommended that the FDA focus on improving its scientific knowledge of nanotechnology to help ensure the agency's regulatory ef-

fectiveness, particularly with regard to products not subject to premarket authorization requirements.

The FDA has already reviewed and approved some nanotechnology-based products. In the coming years, they expect a significant increase in the use of nanoscale materials in drugs, devices, biologics, cosmetics, and food. This will require the FDA to devote more of its regulatory attention to nanotechnology based products.

Let me talk for a few minutes about two areas where nanotechnology is already being applied to health care.

The early detection of cancer and multifunctional therapeutics.

The early detection of cancer can result in significant improvement in human health care and reduction in cost. Nanotechnology offers important new tools for detection where existing and more conventional technologies may be reaching their limits. The present obstacle to early detection of cancer lies in the inability of existing tools to detect these molecular level changes directly during early phases in the genesis of a cancer. Nanotechnology can provide smart contrast agents and tools for real time imaging of a single cell and tissues at the nanoscale.

Nanotechnology promises a host of minimally-invasive diagnostic techniques and much research is aimed at ultra-sensitive labeling and detection technologies. In the *in vitro* area, nanotechnology can help define cancers by molecular signatures denoting processes that reflect fundamental changes in cells and tissues that lead to cancer. Already, investigators have developed novel nanoscale *in vitro* techniques that can analyze genomic variations across different tumor types and distinguish normal from malignant cells.

In the *in vivo* area, one of the most pressing needs in clinical oncology is for imaging agents that can identify tumors that are far smaller than is possible with today's technology. Achieving this level of sensitivity requires better targeting of imaging agents and generation of a larger imaging signal, both of which nanoscale devices are capable of accomplishing.

Perhaps the greatest near-term impact of multifunctional therapeutic compounds will come in the area of tumor targeting and cancer therapies. Nanotechnology can be used to develop new methods of drug delivery that better target selected tissues and cells, and to improve on the efficiency of drug activity in the cytoplasm or nucleus. Drug delivery applications will provide a solution to solubility problems, as well as offer intracellular delivery possibilities.

The introduction of nanotechnology to multifunctional therapeutics is at an early stage of development. The delivery of nanoscale multifunctional therapeutics could permit very precise site specific targeting of cancer cells. More sophisticated "smart" systems for drug delivery still have to be devel-

oped that sense and respond to specific chemical agents and are tailored to each patient. Multifunctional therapeutic devices need to be developed that simultaneously detect, diagnose, treat and monitor response to the therapy. For example, various nanomaterials can be made to link with a drug, a targeting molecule and an imaging agent to seek out cancers and release their payload when required.

The FDA has already begun to devote some resources to the understanding of the human health effects and safety of nanotechnology. It has established a Nanotechnology Core Facility at the FDA's Jefferson Arkansas Laboratories. Combining the expertise of the National Center for Toxicological Research and the Arkansas Research Laboratory, which is part of the FDA Office of Regulatory Affairs, this new Nanotechnology Core Facility will support nanotechnology toxicity studies, develop analytical tools to quantify nanomaterials in complex matrices, and develop procedures for characterizing nanomaterials in FDA-regulated products.

In conclusion, the Nanotechnology Safety Act of 2010 will provide the FDA the authority necessary to scientifically study the safety and effectiveness of nanotechnology-based drugs, delivery systems, medical devices, orthopedic implants, cosmetics, and food additives regulated by the agency. This bill is a sound investment on the promise of nanotechnology to improve human health and reduce costs in the 21st Century.

Mr. President, I ask unanimous consent that the text of the bill be printed in the RECORD.

There being no objection, the text of the bill was ordered to be printed in the RECORD, as follows:

S. 2942

Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

SECTION 1. SHORT TITLE.

This Act may be cited as the "Nanotechnology Safety Act of 2010".

SEC. 2. NANOTECHNOLOGY PROGRAM.

Chapter X of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 391 et seq.) is amended by adding at the end the following:

"SEC. 1011. NANOTECHNOLOGY PROGRAM.

"(a) IN GENERAL.—Not later than 180 days after the date of enactment of the Nanotechnology Safety Act of 2010, the Secretary of Health and Human Services, in consultation with the Secretary of Agriculture, shall establish within the Food and Drug Administration a program for the scientific investigation of nanoscale materials included or intended for inclusion in FDA-regulated products, to address the potential toxicology of such materials, the effects of such materials on biological systems, and interaction of such materials with biological systems.

"(b) PROGRAM PURPOSES.—The purposes of the program established under subsection (a) shall be to—

"(1) assess scientific literature and data on general nanoscale material interactions with biological systems and on specific nanoscale materials of concern to Food and Drug Administration;

"(2) develop and organize information using databases and models that will enable

the formulation of generalized principles for the behavior of classes of nanoscale materials with biological systems;

“(3) promote intramural Administration programs and participate in collaborative efforts, to further the understanding of the science of novel properties at the nanoscale that might contribute to toxicity;

“(4) promote and participate in collaborative efforts to further the understanding of measurement and detection methods for nanoscale materials;

“(5) collect, synthesize, interpret, and disseminate scientific information and data related to the interactions of nanoscale materials with biological systems;

“(6) build scientific expertise on nanoscale materials within such Administration;

“(7) ensure ongoing training, as well as dissemination of new information within the centers of such Administration, and more broadly across such Administration, to ensure timely, informed consideration of the most current science;

“(8) encourage such Administration to participate in international and national consensus standards activities; and

“(9) carry out other activities that the Secretary determines are necessary and consistent with the purposes described in paragraphs (1) through (8).

“(c) PROGRAM ADMINISTRATION.—

“(1) PROGRAM MANAGER.—In carrying out the program under this section, the Secretary shall designate a program manager who shall supervise the planning, management, and coordination of the program.

“(2) DUTIES.—The program manager shall—

“(A) develop a detailed strategic plan for achieving specific short- and long-term technical goals for the program;

“(B) coordinate and integrate the strategic plan with investments by the Food and Drug Administration and other departments and agencies participating in the National Nanotechnology Initiative; and

“(C) develop intramural Administration programs, contracts, memoranda of agreement, joint funding agreements, and other cooperative arrangements necessary for meeting the long-term challenges and achieving the specific technical goals of the program.

“(d) REPORTS.—Not later than March 1, 2012 and March 1, 2014, the Secretary shall submit to the Committee on Health, Education, Labor, and Pensions and the Committee on Appropriations of the Senate and the Committee on Energy and Commerce and the Committee on Appropriations of the House of Representatives a report on the program carried out under this section. Such report shall include—

“(1) a review of the specific short- and long-term goals of the program;

“(2) an assessment of current and proposed funding levels for the program, including an assessment of the adequacy of such funding levels to support program activities; and

“(3) a review of the coordination of activities under the program with other departments and agencies participating in the National Nanotechnology Initiative.

“(e) AUTHORIZATION OF APPROPRIATIONS.—There are authorized to be appropriated to carry out this section, \$25,000,000 for each of fiscal years 2011 through 2015. Amounts appropriated pursuant to this subsection shall remain available until expended.”.

By Ms. COLLINS (for herself, Mr. LIEBERMAN, Mr. BENNETT, Mr. ENSIGN, and Mr. BOND):

S. 2943. A bill to require the Attorney General to consult with appropriate officials within the executive branch

prior to making the decision to try an unprivileged enemy belligerent in Federal civilian court; to the Committee on the Judiciary.

Ms. COLLINS. Mr. President, yesterday the Senate Homeland Security Committee heard testimony from the three top U.S. intelligence officials about the errors that the Federal Government made leading up to the thwarted Christmas Day plot. We dodged a bullet that day when Umar Farouk Abdulmutallab, a Nigerian-born terrorist, failed to detonate a bomb on flight 253 in the skies above Detroit.

But today, Mr. President, I rise to discuss an error that was made after that foreign terrorist had already been detained by American authorities in Detroit, an error that may well have prevented the collection of valuable intelligence about future terrorist threats to our country. The error became clear during my questioning of three of our Nation's top intelligence officials at the committee's hearing yesterday. Frankly, Mr. President, I was stunned to learn that the decision to place the captured terrorist into the U.S. civilian criminal court system had been made without any input or the knowledge of the Director of National Intelligence, the Director of the National Counterterrorism Center, or the Secretary of the Department of Homeland Security. That is right, Mr. President, these officials were never consulted by the Department of Justice before the decision was made.

That decision was critical. The determination to charge Abdulmutallab in civilian court likely foreclosed the collection of additional intelligence information. We know that the interrogation of terrorists can provide critical intelligence, but our civil justice system, as opposed to the military detention and tribunal system established by Congress and the President, encourages terrorists to lawyer up and to stop answering questions. Indeed, that was exactly what happened in the case of Abdulmutallab. He had provided some valuable information to law enforcement officials in the hours immediately after his capture, and we surely would have obtained more information if we had treated this foreign terrorist as an enemy belligerent and had placed him in the military tribunal system. Instead, once he was read his Miranda rights, given a lawyer at our expense, he was advised to cease answering questions, and that is exactly what he did.

That poor decisionmaking may well have prevented us from finding out more of Yemen's role in training terrorists and more about future plots that are underway in Yemen targeting American citizens in this country or abroad. Good intelligence is clearly critical to our ability to stop terrorist plots before they are executed. We know that lawful interrogations of terrorist suspects can provide important intelligence. To charge Abdulmutallab

in the civilian criminal system without even consulting three of our Nation's top intelligence officials simply defies common sense.

To correct this failure and to ensure that our Nation's senior intelligence officials are consulted before making the decision to try future foreign terrorists in civilian court, I am today introducing a bill that would require this crucial consultation. I am very pleased to be joined by the chairman of the Homeland Security Committee, Senator LIEBERMAN, who has been such a leader in this entire area, as well as by three other Senators, Senator BOB BENNETT, Senator JOHN ENSIGN and Senator KIT BOND, who are also concerned about the testimony yesterday.

Specifically, our bill would require the Attorney General to consult with the Director of National Intelligence, the Director of the National Counterterrorism Center, the Secretary of Homeland Security, and the Secretary of Defense before initiating a custodial interrogation of foreign terrorists or filing civilian criminal charges against them. These officials, Mr. President, are in the best position to know what other threats the United States is facing from terrorists and to assess the need to gather more intelligence on those threats.

If there is a disagreement between the Attorney General and these intelligence officials regarding the appropriate approach to the detention and interrogation of foreign terrorists, then the bill would require the President to resolve the disagreement. Only the President would be permitted to direct the initiation of civilian law enforcement actions—balancing his constitutional responsibilities as Commander in Chief and as the Nation's chief law enforcement officer.

To be clear, this legislation would not deprive the President of any investigative or prosecutorial tool. It would not preclude a decision to charge a foreign terrorist in our military tribunal system or in our civilian criminal justice system. It would simply require that the Attorney General coordinate and consult with our top intelligence officials before making a decision that could foreclose the collection of critical additional intelligence information.

This consultation requirement is not unprecedented. Section 811 of the Counterintelligence and Security and Enhancements Act of 1994 requires the Director of the FBI and the head of a department or agency with a potential spy in its ranks to consult and periodically reassess any decision to leave the suspected spy in place so that additional intelligence can be gathered on his activities.

As the Senate Intelligence Committee noted in its report on the legislation that added the espionage consultation requirement:

While prosecutorial discretion ultimately rests with the Department of Justice officials, it stands to reason that in cases designed to protect our national security—such

as espionage and terrorism cases—prosecutors should ensure that they do not make decisions that, in fact, end up harming the national security.

The committee got it right. The committee went on to explain:

[T]he determination of whether to leave a subject in place should be retained by the host agency.

The history of the espionage consultation requirement is eerily reminiscent of the lack of consultation that occurred in the case of Abdulmutallab. In espionage cases, Congress has already recognized that when valuable intelligence is at stake, our national security should trump decisions based solely on prosecutorial equities. This requirement must be extended to the most significant threat facing our Nation, and that is the threat of terrorism.

I encourage the Senate to act quickly on this important legislation. The changes proposed are modest. They make common sense. But the consequences could be a matter of life and death.

Mr. President, I ask unanimous consent that the text of the bill be printed in the RECORD.

There being no objection, the text of the bill was ordered to be printed in the RECORD, as follows:

S. 2943

Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

SECTION 1. CONSULTATION REQUIREMENT.

(a) IN GENERAL.—Subject to subsection (b), no action shall be taken by the Attorney General, or any officer or employee of the Department of Justice, to—

- (1) initiate a custodial interrogation of; or
- (2) file a civilian criminal complaint, information, or indictment against;

any foreign person detained by the United States Government because they may have engaged in conduct constituting an act of war against the United States, terrorism, or material support to terrorists, or activities in preparation therefor.

(b) CONSULTATION.—

(1) IN GENERAL.—Subject to paragraph (2), the Attorney General shall consult with the Director of National Intelligence, the Director of the National Counterterrorism Center, the Secretary of Homeland Security, and the Secretary of Defense prior to taking any action identified in subsection (a).

(2) PRESIDENTIAL DIRECTION.—If, following consultation under paragraph (1), the Director of National Intelligence, the Director of the National Counterterrorism Center, the Secretary of Homeland Security, or the Secretary of Defense believe that any action identified in subsection (a) and proposed by the Attorney General may prevent the collection of intelligence related to terrorism or threats of violence against the United States or its citizens, the Attorney General may not initiate such action without specific direction from the President.

(c) ANNUAL REPORT.—The Attorney General shall report annually to appropriate committees of jurisdiction regarding the number of occasions on which direction was

sought from the President under subsection (b)(2) and the number of times, on those occasions, that the President directed actions identified in section (a) against such foreign person.

(d) DEFINITIONS.—In this section:

(1) APPROPRIATE COMMITTEES OF JURISDICTION.—The term “appropriate committees of jurisdiction” shall include—

(A) the Committee on Homeland Security and Governmental Affairs of the Senate;

(B) the Committee on Homeland Security of the House of Representatives;

(C) the Select Committee on Intelligence of the Senate;

(D) the Permanent Select Committee on Intelligence of the House of Representatives; and

(E) the Committees on Armed Services and Judiciary of the Senate and the Committees on Armed Services and Judiciary of the House of Representatives.

(2) ACT OF WAR, TERRORISM, MATERIAL SUPPORT TO TERRORISTS.—The terms “act of war”, “terrorism”, and “material support to terrorists” shall have the meanings given such terms in title 18, United States Code.

(e) SAVINGS CLAUSE.—Nothing in this section shall prevent the Attorney General, or any officer or employee of the Department of Justice, from apprehending or detaining an individual as authorized by the Constitution or laws of the United States except to the extent that activities incident to such apprehension or detention are specifically identified in subsection (a).

SUBMITTED RESOLUTIONS

SENATE RESOLUTION 390—PROHIBITING TEXT MESSAGING BY EMPLOYEES OF THE SENATE WHILE DRIVING ON OFFICIAL BUSINESS

Mr. CASEY (for himself, Mr. SCHUMER, and Mr. ROCKEFELLER) submitted the following resolution; which was referred to the Committee on Rules and Administration:

S. RES. 390

Resolved,

SECTION 1. PROHIBITION ON TEXT MESSAGING BY EMPLOYEES OF THE SENATE WHILE DRIVING ON OFFICIAL BUSINESS.

(a) DEFINITIONS.—In this resolution—

(1) the term “employee of the Senate” means any employee whose pay is disbursed by the Secretary of the Senate; and

(2) the term “text messaging” means reading from or entering data into any handheld or other electronic device, including for the purpose of SMS texting, e-mailing, instant messaging, obtaining navigational information, or engaging in any other form of electronic data retrieval or electronic data communication.

(b) PROHIBITION.—An employee of the Senate may not engage in text messaging when—

(1) driving a Government owned or leased vehicle;

(2) driving a privately owned or leased vehicle while on official business; or

(3) using text messaging equipment provided by any officer or committee of the Senate while driving any vehicle at any time.

(c) EFFECTIVE DATE AND APPLICATION.—This resolution shall apply to the 111th Congress and each Congress thereafter.

SENATE RESOLUTION 391—RECOGNIZING THE 25TH ANNIVERSARY OF THE ENACTMENT OF THE VICTIMS OF CRIME ACT OF 1984 (42 U.S.C. 10601 ET SEQ.) AND THE SUBSTANTIAL CONTRIBUTIONS TO THE CRIME VICTIMS FUND MADE THROUGH THE CRIMINAL PROSECUTIONS CONDUCTED BY UNITED STATES ATTORNEYS' OFFICES AND OTHER COMPONENTS OF THE DEPARTMENT OF JUSTICE

Mr. CRAPO (for himself, Ms. KLOBUCHAR, and Mr. VITTER) submitted the following resolution; which was considered and agreed to:

S. RES. 391

Whereas the Victims of Crime Act of 1984 had its 25th anniversary in 2009;

Whereas for 25 years, the Victims of Crime Act of 1984 has provided funds to States for victim assistance and compensation programs to support victims of crime and those affected by violent crimes;

Whereas the Victims of Crime Act of 1984 enables approximately 4,400 community-based public and private programs to offer services to victims of crime, including crisis intervention, counseling, guidance, legal advocacy, and transportation shelters;

Whereas the Victims of Crime Act of 1984 provides assistance and monetary support to over 4,000,000 victims of crime each year;

Whereas the Crime Victims Fund established under the Victims of Crime Act of 1984 provides direct services to victims of sexual assault, domestic violence, child abuse, survivors of homicide victims, elderly victims of abuse or neglect, victims of drunk drivers, and other such crimes;

Whereas in 2008, with financial support from the Victims of Crime Act of 1984, State crime victim compensation programs paid a total of \$432,000,000 to 151,643 victims of violent crime;

Whereas since the establishment of the Crime Victims Fund in 1984, non-taxpayer offender-generated funds deposited into the Crime Victims Fund have been used to provide almost \$7,500,000,000 to State crime victim assistance programs and State crime victim compensation programs;

Whereas the Victims of Crime Act of 1984 also supports services to victims of Federal crimes, by providing funds for victims and witness coordinators in United States Attorneys' offices, Federal Bureau of Investigation victim-assistance specialists, and the Federal Victim Notification System; and

Whereas the Victims of Crime Act of 1984 also supports important improvements in the victim services field through grants for training and technical assistance and evidence-based demonstration projects: Now, therefore, be it

Resolved, That the Senate recognizes—

(1) the 25th anniversary of the enactment of the Victims of Crime Act of 1984 (42 U.S.C. 10601 et seq.); and

(2) the substantial contributions to the Crime Victims Fund made through the criminal prosecutions conducted by United States Attorneys' offices and other components of the Department of Justice.