spinal nerves both come through an open part of the spine.

While medical science has not vet discovered the exact cause of Spina Bifida-it is a birth defect that can strike anyone-we do know that a woman taking 400 mcg of folic acid whether as part of a multivitamin/drug formula or alone, prior to and in the early weeks of pregnancy, reduces her risk of having a baby with Spina Bifida or another neural tube defect by as much as 70 percent. That is why since February 1996 and at the direction of Food and Drug Administration, all U.S. food manufacturers started adding folic acid to most enriched breads, flours, corn meals, pastas, rice and other grain products to reduce the risk of birth defects in newborns. Following the fortification of cereal grains, the incidences of these types of birth defects decreased by about 20 percent.

However, surveys and research suggest that only about 30 percent of American women consume the daily recommended amount of folic acid. As a result, today in America there are still some 60 million women at risk of having a baby born with Spina Bifida, and everyday, an average of 8 babies are affected by Spina Bifida or a similar birth defect of the brain and spine. Annually, approximately 3,000 pregnancies are affected by these birth defects.

These devastating birth defects cannot be cured, so they must be prevented, and the evidence of folic acid's benefits as a preventative is extremely compelling. The solution is education and awareness. Since 1973, the Spina Bifida Association of America—the nation's only organization solely dedicated to advocating on behalf of the Spina Bifida community-has been leading the charge. Through their almost 60 chapters in more than 125 communities, the SBAA has helped bring thousands of expectant parents together with parents of children who suffer from the disease. This type of one-on-one interaction helps concerned parents get answers to their questions, but most importantly it lends support and gives people essential information and hope. Thanks to SBAA's efforts, millions of people have received vital information about Spina Bifida and how it affects those who live with it; countless women have learned about the importance of taking folic acid prior to pregnancy to reduce the risk of Spina Bifida and other neural tube defects; and, countless individuals afflicted with Spina Bifida have gotten the help they need to live life to its fullest and achieve their full potential.

A very special lady who works with me in my office, Ms. Donna Jones, in fact happens to have Spina Bifida. Since the day she came on board, she has done a great job and helped me understand first-hand the challenge of living with this disease. Ever since I've known her, Donna has always believed that change can come if even one individual believes in themselves and strives to make a difference no matter how hard the challenge. And she has never let the challenge of living with her disease slow her down or stop her from pursuing her dream of making a difference and advocating for persons with disabilities. I was happy to be able to give her the opportunity to come to Washington and try to make her dream a reality; and I have no doubt that because of her tireless efforts to raise awareness of Spina Bifida on Capitol Hill, the incidence of Spina Bifida will decrease and fewer children will have to endure the road she has had to travel.

Even with all the challenges she faces, I have never seen Donna without a smile on her face. She definitely has an infectious love of life, and when the world looks at her it doesn't see a person unobtrusively going through the motions of everyday life, it sees an outspoken, dynamic and active person willing to get involved in the issues that she cares about, particularly those that affect disabled individuals in our country.

I believe that Donna is perhaps the best advocate the Spina Bifida community could have on Capitol Hill because each and every day, she serves as living proof that someone with Spina Bifida need never take a back seat to anyone. I wish more young people in America had the same fire and commitment as Donna. In fact, it is largely thanks to Donna that I became aware of and joined the Congressional Spina Bifida Caucus. I hope that many of my colleagues listening today will also join the Caucus and work to spread awareness and education about this disease. Working together we can do so much to improve the quality of life to those living with Spina Bifida.

Just this past Wednesday, October 5, 2005, the SBAA's held its 17th Annual Gala here in Washington to benefit the Association and its many chapters around the nation. The event was a tremendous success and the funds raised will help the SBAA and its chapters around the country continue the good work they do in preventing more occurrences of this devastating disease, and reducing the suffering of those 70,000 Americans living with Spina Bifida. I firmly believe that we owe SBAA a great debt for what they have accomplished.

In my home State of Indiana, SBAA, the Spina Bifida Association (SBA) of Central Indiana and the SBA of Northern Indiana have formed a collaborative partnership to make a difference in the lives of Hoosier families living with Spina Bifida by helping them meet the challenges and enjoy the rewards of raising their children. In addition, in response to the devastation of the Gulf region of this country by Hurricanes Katrina and Rita, Indiana's Spina Bifida chapters answered the call by participating in the Spina Bifida Hurricane Emergency Life Support Program (SBHELSP). Established by the SBAA, the SBHELSP, assisted in getting vitally needed supplies to victims of the hurricanes and to put displaced families affected by Spina Bifida in touch with others who understand their special needs and can help them. I cannot say enough good things about the hard work, dedication and compassion of the volunteers and staff at all the Indiana chapters of SBAA: I can only thank them on behalf of all Hoosiers for all they have done and all that they will continue to do.

SBAA and its local chapters are making a difference one life at a time. With National Spina Bifida Awareness Month upon us again, I would say to my colleagues in closing that the time for us to start making a difference is now

CONTINUING APPROPRIATIONS, FISCAL YEAR 2006

SPEECH OF

HON. JAMES R. LANGEVIN

OF RHODE ISLAND

IN THE HOUSE OF REPRESENTATIVES
Thursday, September 29, 2005

Mr. LANGEVIN. Mr. Speaker, I rise in opposition to H.J. Res. 68, the Continuing Appropriations Resolution for FY 2006. By funding the government at the lower of the House bill, the Senate bill, or current rate, this continuing resolution would require immediate cuts in programs that address the very problems that America's families are facing this fall—unemployment, education, health care, and emergency preparedness.

Right now, there are 7.4 million unemployed Americans—1.4 million of whom have been jobless for more than 26 weeks. Thousands more are applying for benefits in the wake of Katrina and Rita, yet this bill would cut essential Federal training and employment services below last year's level, including \$138 million, or 4 percent, from job training formula grant programs. Just as students are going back to school, this legislation calls for an \$800 million cut in education programs. With 45.8 million uninsured Americans, this bill fails to provide necessary assistance to the Community Health Centers that serve these very citizens. In a time when emergency preparedness should be a high priority, the bill cuts Hospital Preparedness Grants by \$15 million and Public Health Preparedness Grants by \$66 million.

In light of these pressing concerns for all Americans and rising defecits, we must spend our dollars wisely. Yet, we have been presented with a bill that shortchanges the very programs that promise to rebuild our workforce, educate our children and protect the health and well being of our citizens. I urge my colleagues to join me in supporting fiscal responsibility by voting against H.J. Res. 68.

A SPECIAL TRIBUTE TO STARR COMMONWEALTH ON THE OCCASION OF ITS EXPANDED MONTCALM SCHOOL FOR GIRLS

HON. PAUL E. GILLMOR

OF OHIO

IN THE HOUSE OF REPRESENTATIVES

Thursday, October 6, 2005

Mr. GILLMOR. Mr. Speaker, it is my privilege to pay tribute to a special organization in Ohio's Fifth Congressional District. This October, the Montcalm School for Girls at Starr Commonwealth's Van Wert campus celebrates its first anniversary celebration.

Mr. Speaker, Starr Commonwealth has been nationally recognized for serving thousands of children, families and professionals throughout America. For nearly a century, the dedicated staff of Starr Commonwealth has committed themselves to diffusing violence, healing racism and developing positive environments in the children which they serve. With five locations in Michigan and Ohio, Starr Commonwealth operates dozens of programs for troubled youth and their families. By instilling the core values of integrity, excellence, compassion and faith, Starr Commonwealth has been effective at redirecting misdirected children across our nation.

In 1951, Floyd Starr expanded Starr Commonwealth to a campus outside of Van Wert in Ohio's Fifth Congressional District. Staff members at the Van Wert campus help boys with social, emotional and behavioral problems as they attempt to turn their lives around. In addition, the school is celebrating its one-year anniversary of operating the Montcalm School for Girls. The Montcalm School provides residential treatment for young women who have experienced abuse, depression, behavioral or academic problems.

The services provided by Starr Commonwealth and the Montcalm School are good for our children and good for our society. Students who attend the Montcalm School have fallen off the path in life and need our assistance to become productive members of society once again. These students have engaged in activities which warrant our attention but they also possess a passion to be rehabilitated. By attending the Montcalm School, students are provided with a therapeutic environment to allow them to grow as individuals and avoid a life of delinquency. The school boasts a 95 percent success rate and has given back to our society those who had before been forgotten.

Mr. Speaker, it is not everyday that we can recognize programs which provide such an important service to our community as Starr Commonwealth. In Van Wert, citizens of the Fifth District have committed to protecting and rehabilitating our children through the Montcalm School. Mr. Speaker, I ask my colleagues to join me in paying special tribute to Starr Commonwealth for its commitment to a better America. As all who benefit from Starr Commonwealth gather to celebrate the one-year anniversary of the Montcalm School, I am confident that the services provided in Van Wert will continue to better our society into the future.

TRIBUTE TO THE LATE HOWARD BEAVER

HON. RAUL M. GRIJALVA

OF ARIZONA

IN THE HOUSE OF REPRESENTATIVES Thursday, October 6, 2005

Mr. GRIJALVA. Mr. Speaker, it is with regret and deep sadness that I rise to honor Howard Beaver, a resident of Tucson, AZ, who passed away on October 3, 2005. For over 27 years he sold or rented thousands of musical instruments to Tucson children. I myself bought my first harmonica at his store. He was a pillar of Tucson society, an impassioned musician, beloved husband, father, and grandfather.

Howard Beaver was born in Beaver, AR, and raised in Odessa, TX. He moved to Tucson in 1949 for his freshman year at the University of Arizona, where he received his bachelor's degree in music. He later went on to earn a master's degree from Northern Arizona University.

Howard opened Beaver's Band Box in 1964, a music store where he rented and sold instruments as well as provided music lessons. In 1991 he sold the store to his son, Dan, who then proceeded to open a second Beaver's Band Box.

He was a very dynamic member of the Tucson music community; in 1966 he founded Beaver's Tucson Concert Band, was influential in the beginning of the UA alumni band and was active in many community bands.

Howard's life was filled with music and family, two things that were very important to him. His love of music began early in life, with his first Trombone in seventh or eighth grade, though he would later know how to play them all. The other love of his life was family. He is survived by his wife Liz, five children (Dan, Heidi, Carl, Krista, and Erica) and 15 grand-children.

TRIBUTE TO HISPANICS ACHIEVING AND CELEBRATING EXCELLENCE

HON. JAMES P. McGOVERN

OF MASSACHUSETTS

IN THE HOUSE OF REPRESENTATIVES Thursday, October 6, 2005

Mr. McGOVERN. Mr. Speaker, I rise today to recognize the achievements of Ms. Lucia Alfaro, Ms. Cristina Cartagena, Ms. Amanda Ms. Genesis Melendez, Ms. Gonzalez, Daniela Ortiz Bahamonde, Mr. Esteban Pacheco, Ms. Sylvia Roman, Mr. Enrique Sanchez, Ms. Elizabeth Zamora, Mr. David Hernandez, Ms. Mayra Jara, Mr. Josue Malaver, Ms. Sujeilly Rohena, Ms. Linette Serrano, Mr. Nolberto Alvarenga, Mr. Angel Amarat, Mr. Eric Familia, Mr. Karl Gonzalez, Ms. Makisha Rodriguez, Mr. Cruz Rosado, Mr. Benito Vega, Ms. Katherine Cortes, Ms. Carol Esparza, Mr. Narcisco Fernandez, Mr. Miguel Fuentes, Ms. Jessica Perez, Mr. Eric Sanchez, and Ms. Christina Gonzalez. These fine young people from the city of Worcester have recently been recognized at the annual HACE Youth Awards and Recognition Dinner.

The Hispanics Achieving and Celebrating Excellence Committee, HACE, was created in 1985 by the faculty and staff at Quinsigamond Community College. The goal of HACE is to raise community awareness of the richness of Hispanic culture. The organization recognizes the achievements of young people, provides positive role models, and encourages youths to focus on education. Honorees are nominated by school principals, teachers, coaches, community leaders and agencies. Along with the honor, each student receives a scholarship.

Each year, HACE holds an annual dinner to recognize youth leaders. The honorees from this year's annual dinner, include 28 students from Worcester high schools. These select few have distinguished themselves in five categories: the arts, athletics, academics, community service and/or leadership. This year's honorees will receive a gift in the amount of \$200, as an incentive and encouragement to continue along the path of excellence.

Over a span of 20 years, HACE is proud to have given monetary awards totaling over \$57,000 in recognition of the achievements each honoree has accomplished.

The committee members who dedicate themselves to coordinating HACE include Ms. Maria Addison, Ms. Olga Lopez-Hill, Mr. John Rodriguez, Mr. Dan de la Torre, Ms. Susan Laprade, Ms. Brenda Safford, Ms. Kirshner Donis, Ms. Viviana Troche, Honorable Harriette Chandler, Ms. Maria Florez, Mr. Javier Salort, Mr. Albert Vasquez, and Ms. Dolly Vazquez.

I am sure all my colleagues and fellow constituents will join me in applauding the accomplishments of these young leaders, and praise HACE for the great service they perform in the city of Worcester.

TRIBUTE TO SERGEANT ANDREW JOSEPH DERRICK

HON. JOE WILSON

OF SOUTH CAROLINA

IN THE HOUSE OF REPRESENTATIVES

Thursday, October 6, 2005

Mr. WILSON of South Carolina. Mr. Speaker, Sgt. Andrew Joseph Derrick of Columbia, South Carolina died in Baghdad, Iraq on September 25, 2005. I am greatly saddened by his death and humbled by his sacrifice. As citizens throughout our community continue to cope with this tragic loss, my thoughts and prayers will remain with Sgt. Derrick's family, his friends and his fellow service members.

The following obituary was published in The State newspaper on Saturday, October 1, 2005.

Joseph was a loving son, brother and fiance. Known for his positive attitude and playfulness, he was able to uplift the spirits of everyone he encountered. He was a superlative soldier and a man of honor and integrity. As a man of strong Christian faith, he was selfless, giving, kind, tenderhearted and never afraid to share the hope he had in Christ. He now rests in eternal peace with his Savior.

Sgt. Derrick died Sunday, September 25, 2005, while stationed in Baghdad, Iraq. Born in Columbia, he was a son of Suzanne McCormick Shealy and Andrew Carroll "Butch" Derrick. He was a 1998 graduate of Marion Military Institute and attended Francis Marion University. While a student at Marion Military Institute, Joseph played and lettered in Varsity football, baseball and basketball. He was named the Class of 1998's "Best All Round Athlete" and was named to the State of Alabama's All Star Baseball Team in 1998.

During his military career, Joseph graduated from the Primary Leadership Development Course and was awarded the Bronze Star, the Purple Heart, the Army Achievement Medal (with one Oak Leaf Cluster), the National Defense Service Ribbon, the Iraqi Campaign Medal, the Global War on Terror Medal, the Korean Defense Service medal, the Army Service Ribbon, the Combat Action Badge, the Marksmanship Badge, the Good Conduct Medal, and the Army Commendation Medal.

Surviving are his mother, Suzanne "Suzv" McCormick Shealy of Columbia: stepfather. Carey William Shealy of Columbia; brother, William James Shealy of Columbia; sister, Elizabeth Anne Shealy of Columbia: father. Butch Derrick of Columbia and his wife, Deborah R. Derrick; half sister, Chelsea Campbell Derrick and Austin Reid Royal: paternal grandmother, Barbara Jean Shealy of Columbia; maternal aunt and uncle, Mr. and Mrs. L. Stephen Quatannens (Jo Anne) of Campobello, S.C.; Shealy paternal aunt and uncle, Mr. and Mrs. Whitfield Milling Howard (Rebecca) of Surfside Beach, S.C.; paternal uncle, Robert William Derrick of Columbia; paternal aunt, Barbara Rainey of Greensboro, Ga.; first cousins, Mrs. Jamey Scheuch (Andrea), Mrs. Bob Elder (Catherine) and Mr. Kenneth Whitfield Howard; and numerous second cousins. He is also survived by his fiancee, Miss Shaunna Marie Miller of Phoenix, Ariz.

He was predeceased by his uncle, Kenneth James Shealy; grandparents, Samuel James