



## State of Tennessee

### SENATE JOINT RESOLUTION NO. 122

By Yager

A RESOLUTION to designate February 28, 2011, as Rare Disease Day in Tennessee.

WHEREAS, in an effort to raise awareness in the United States about rare diseases, Rare Disease Day, on February 28, 2011, has been designated as the occasion to bring forth a national audience in an attempt to raise cognizance about issues surrounding rare diseases; and

WHEREAS, there are nearly 7,000 diseases and conditions considered rare, each affecting fewer than 200,000 Americans; and

WHEREAS, while each of these diseases may affect small numbers of people, rare diseases as a group affect almost thirty million Americans; and

WHEREAS, many rare diseases are serious and debilitating conditions that have a significant impact on the lives of those affected; and

WHEREAS, while nearly 330 orphan drugs and biologics have been approved for the treatment of rare diseases affecting between eleven and fourteen million people according to the Food and Drug Administration, that leaves well over fifteen million Americans with rare diseases who have no treatment specific to their disease; and

WHEREAS, individuals and families affected by rare diseases often experience problems such as a sense of isolation, difficulty in obtaining an accurate and timely diagnosis, few treatment options and problems related to accessing or being reimbursed for treatment; and

WHEREAS, some rare diseases, such as "Lou Gehrig's Disease" and "Huntington's Disease", are relatively well known; many others are not known at all by the public, so that patients and their families must bear a large share of the burden for things such as raising funds for research; and

WHEREAS, the National Organization for Rare Disorders (NORD) organizes a nationwide observance of Rare Disease Day on February 28th, and patients, medical professionals, researchers, government officials and companies developing treatments for rare diseases join together to focus attention on rare diseases as a public health issue on that day; and

WHEREAS, successes of the annual Rare Disease Day USA include governors in forty states issuing proclamations to recognize the important issue, rarediseaseday.us being launched, more than 350 groups signing on as Rare Disease Day Partners, and patient stories appeared on ABC News, blogs, editorials, newspapers and many other Web sites to address the travailing issue; and

WHEREAS, thousands of residents of our great State are among those affected by rare diseases since nearly one in ten Americans have rare diseases; now, therefore,

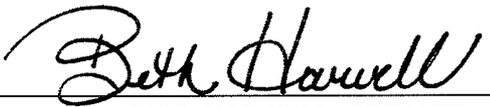
BE IT RESOLVED BY THE SENATE OF THE ONE HUNDRED SEVENTH GENERAL ASSEMBLY OF THE STATE OF TENNESSEE, THE HOUSE OF REPRESENTATIVES CONCURRING, that we join in the designation of February 28, 2011, as Rare Disease Day, and encourage all Tennesseans to educate themselves about the reality of rare diseases across our State and Nation.

BE IT FURTHER RESOLVED, that an appropriate copy of this resolution be prepared for presentation with this final clause omitted from such copy.

SENATE JOINT RESOLUTION NO. 122

ADOPTED: May 16, 2011

  
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RON RAMSEY  
SPEAKER OF THE SENATE

  
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BETH HARWELL, SPEAKER  
HOUSE OF REPRESENTATIVES

APPROVED this 24<sup>th</sup> day of May 2011

  
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BILL HASLAM, GOVERNOR