March 23, 2021

The Honorable William Cloud Robinson, Jr.
Chair
House Professions and Public Health Subcommittee
214 House Building
402 South Monroe Street
Tallahassee, FL 32399

Re: Support for House Bill 1373

Dear Chairman Robinson and Members of the House Professions and Public Health Subcommittee:

On behalf of the 1-in-10 individuals in Florida with one of the approximately 7,000 known rare diseases, the National Organization for Rare Disorders (NORD) thanks you for putting House Bill 1373 (HB 1373) on the House Professions and Public Health Subcommittee’s agenda. HB 1373 establishes a Rare Disease Advisory Council (RDAC), which if passed, would help to give a voice to the rare disease community within Florida’s state government. NORD urges you to vote in support of HB 1373 and swiftly move it out of your Subcommittee.

NORD is a unique federation of voluntary health organizations dedicated to helping people with rare "orphan" diseases and assisting the organizations that serve them. We are committed to the identification, treatment, and cure of rare disorders through programs of education, advocacy, research, and patient services.

Rare diseases are present across a broad spectrum of medical conditions. For example, there are more than 500 types of rare cancers and all forms of pediatric cancer are rare. For patients suffering from these and numerous other rare conditions, it can take several years to receive an accurate diagnosis and effective treatment. Further, only a handful of rare diseases are well-understood, with most not receiving sufficient attention or funding for research.

Creating an RDAC in Florida will give rare disease patients a unified voice in Florida’s state government, helping to provide educational awareness and make recommendations to elected officials and other state leaders on how rare diseases are handled and treated in the state. From providing information on the diagnostic journey, to making recommendations on state programs such as newborn screening, the council will serve as a tremendous opportunity for important decision-makers in Florida to better understand and meet the needs of their constituents. Since this council would include broad participation from the different health care sectors present in Florida, it will also serve as an educational resource to all stakeholders about the ways rare disease patients interact with our health care system.
In creating this council, Florida will be joining sixteen other states (Alabama, Connecticut, Illinois, Kentucky, Massachusetts, Minnesota, Missouri, New Hampshire, New York, Nevada, North Carolina, Ohio, Pennsylvania, Tennessee, Utah, and West Virginia) who have already enacted similar legislation in support of their rare disease community.

Once again, on behalf of the Florida rare disease community, we thank you for putting HB 1373 on the House Professions and Public Health Subcommittee’s agenda and urge its swift passage. For any questions, please feel free to contact Annissa Reed via email at areed@rarediseases.org. Thank you for your consideration.

Sincerely,

Heidi Ross  
Director of Policy  
National Organization for Rare Disorders

Annissa Reed  
State Policy Manager, Eastern Region  
National Organization for Rare Disorders

CC: Members of the House Professions and Public Health Subcommittee
March 23, 2021

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Re: Support for House Bill 1373

Dear Chairman Robinson and Members of the House Professions and Public Health Subcommittee:

My name is Anne Rugari, and I am the Florida Rare Action Network Volunteer State Co-Ambassador for the National Organization for Rare Disorders. NORD is a unique federation of voluntary health organizations dedicated to helping the 25-30 million Americans living with a rare disease.

As a rare disease advocate, parent, and caregiver, I have had many opportunities to not only work with the rare disease community, but to also hear the voices of many people who are affected with a rare disorder or disease. It is estimated that One in 10 Floridians is affected by a rare disorder. A Rare Disease Advisory Council (RDAC) in the State of Florida would address various issues that the rare disease community in Florida experiences, and will work with the Florida Legislature to help improve policies would support the rare disease population in having accessibility to better health care.

My first experience with rare diseases came from having two children born with a rare, genetic, neurologic disorder called Krabbe disease. Krabbe is a demyelinating disease that affects the central and peripheral nervous systems. Children born with this disorder typically die within the first two years of life. My son, Nicholas lived to be a year old. My daughter, Gina, was tested at birth for Krabbe and was able to receive a treatment that helped her to live a much longer and better quality of life than her brother. The many healthcare issues associated with Krabbe, has given me firsthand knowledge of the challenges faced by those in the rare disease community and has prepared me to be a voice and advocate for all of the rare disease community.

It has become increasingly complicated for the rare disease community to receive the support services, prescription drugs, and other medical treatments, even more so during the COVID-19 pandemic. The RDAC would work to alleviate some of these healthcare burdens for this vulnerable population.
To date, sixteen states (Alabama, Connecticut, Illinois, Kentucky, Massachusetts, Minnesota, Missouri, New Hampshire, New York, Nevada, North Carolina, Ohio, Pennsylvania, Tennessee, Utah, and West Virginia) have created a Rare Disease Advisory Council and proven that it is an effective way to ensure better government engagement and action on issues of importance to the rare disease community. I urge you to vote in support of House Bill 1373 and help Florida become the 17th state to enact an RDAC.

Sincerely,

Anne Rugari
Florida Volunteer State Ambassador
Florida Rare Action Network

CC: Members of the House Professions and Public Health Subcommittee
March 23, 2021

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Re: Support for House Bill 1373

Dear Chairman Robinson and Members of the House Professions and Public Health Subcommittee:

My name is Jhoanny Cardenas, and I am the Florida Rare Action Network Volunteer State Co-Ambassador for the National Organization for Rare Disorders. NORD is a unique federation of voluntary health organizations dedicated to helping 25-30 million American living with a rare disease.

As my younger son, Alexander, entered into the first grade, his teachers brought up some behavior changes that they had observed and concerned them.

I began taking my son Alex to the pediatrician to discuss their concerns, which resulted in getting referrals to see other specialists. I was frequently told by these specialists that “there was nothing wrong with your son, he is being a regular six-year-old,” but I was not satisfied with this response. I made additional appointments with a child neurologist and endocrinologist. I was discouraged from doing this because I was told my insurance may not pay as there was no family history, even though I had previously expressed some concerns. It was after several tests ordered by the endocrinologist that on March of 2015 Alexander was diagnosed with Adrenoleukodystrophy (X-ALD). This was a new and scary diagnosis. We understood that time was not on our side. We quickly traveled to Minnesota, where Alexander received treatment.

I began having medical issues in middle school, but I wasn’t diagnosed with Ehlers-Danlos Syndrome until I was 19. My personal diagnostic odyssey took place over seven years and brought me to five different states and almost 15 physicians before I got an accurate diagnosis.

Sadly, Alexander’s diagnosis was not my first connection to the rare disease community, which began at a young age when my younger sister, Luzel, was paralyzed in what we have now been told was due to Gillian-Barre Syndrome. She made a full recovery and lived a “normal” life until seven years ago, when she began to experience debilitating headaches and muscle weakness which at times resembled a stroke. I have supported my sister through research and visiting various specialists throughout the state. Two years ago, she was diagnosed with Dystonia and most recently, Stiff Person Syndrome. I think about the challenges that my sister faces each day with her conditions, and how it impacts her life. I also think about the difference it would have made for our family as we navigated the new diagnosis and all that came along with Alexander’s...
care. Alexander went from being a healthy and active six-year-old, to very medically complex. Had he been diagnosed earlier; his story may have had a different ending. Just a short seventeen months after diagnosis, Alexander experienced complications after two transplants which he was not able to overcome and lost his battle to Adrenoleukodystrophy in October of 2016, at just eight years old.

Navigating any medical diagnosis can be challenging and complicated and a rare disorder add even more complexity. Many of us find ourselves not knowing where to turn and feel alone and without a voice. A RDAC can help guide and encourage all stakeholders to collaborate and help provide recommendations to improve the diagnosis, care and treatment of rare disease patients. I urge you to vote in support of House Bill 1373 and help Florida become the 17th state to enact an RDAC.

Sincerely,

Jhoanny Cardenas
Florida Volunteer State Ambassador
Florida Rare Action Network

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