Dear Members of Congress:

As an organization focused on rare disease, I am signing on to support appropriations for Fiscal Year 2022 Labor, Health, Human Services, Education and Related Agencies to ensure every state has fully implemented all Recommended Uniform Screening Panel (RUSP) conditions.

While the federal government makes recommendations on diseases for newborn screening, it is implemented on a state by state basis. Every year, thousands of children die or become permanently disabled simply because they are not screened at birth for many known genetic diseases that have a treatment, resulting in something known as "Death by Zip Code" -- infants in one state are identified with a debilitating disease but those in another are not. This doesn't have to happen.

In 2019 the Secretary of HHS's Advisory Committee on Heritable Disorders in Newborns and Children received the results of the New STEPs New Disorders study which identified the two greatest barriers to implementation of RUSP newborn screening to be state staffing needs and equipment procurement. Currently, only 15 states are implementing newborn screening for all RUSP conditions. A number of states have not yet begun to screen newborns for these rare diseases even though in several instances they were added by the RUSP more than 6 years ago.

As we move forward at the appropriate time with the Fiscal Year 2022 Labor, Health, Human Services, Education and Related Agencies bill, we urge you to ensure every state has fully implemented newborn screening for all Recommended Uniform Screening Panel (RUSP) conditions. Specifically, we ask you to direct the Centers for Disease Control (CDC) to provide \$15 million for timely implementation of newborn screening conditions with a goal of complete RUSP implementation in all 50 states by 2025.

At present rates, it will take states more than a decade to implement newborn screening for RUSP approved diseases for which treatment options are available to families, resulting in preventable deaths and disability. The Centers for Disease Control and Prevention has existing legislative authority and direct experience working with states to implement RUSP recommendations for enhanced newborn screening. The House of Representatives also recently demonstrated strong, bipartisan support for advancing newborn screening with the passage of H.R. 2507, the Newborn Screening Saves Lives Reauthorization Act of 2019.

In closing, we request that you ensure that children in all 50 U.S. states have a fair chance at life. We know from experience that early diagnosis will save countless children from death or paralysis. This is a life-and-death issue. To end "Death by Zip Code," we must prioritize complete RUSP implementation by all 50 states by 2025. Accordingly, we encourage you to include both funding and report language in the Fiscal Year 2022 Labor HHS appropriations bill mandating such implementation. With your leadership and support, we can save lives and improve the health of newborns in all 50 U.S. states.

Sincerely,

Acid Maltase Deficiency Association

Adrenal Insufficiency United Aidan Jack Seeger Foundation

Alabama Rare ALD Alliance ALD Connect

Alexander Matthew Foundation

Arrivederci ALD

Association for Creatine Deficiencies

Believing for Bryleigh

Benji Strong

Beyond the Diagnosis Organization

Brian's Hope

Bridge the Gap - SYNGAP Education and

Research

Calliope Joy Foundation

Center for Independence of the Disabled NY

Children's Hospital of Pennsylvania Children's Healthcare of Atlanta

Cure AHC

Friedreich's Ataxia Research Alliance (FARA)

Cure San Filippo Cure VCP Disease

Cutaneous Lymphoma Foundation

Dalton's New World Order Dysautonomia Determination Ehlers Danos Hypermobility Ethan Zakes Foundation

Fight ALD

Foundation to Fight H-abc Gene Giraffe Project Gene Spotlight Charity

Global Foundation Peroxisomal Diseases

Global Genes

Global Leukodystrophy Initiative

Hope for Dante Hunter's Hope Hydrocephalus Kids

Icahn School of Medicine at Mount Sinai

Jonah's Just Begun Judson's Legacy

Kennedy Krieger Institute

Krabbe Connect

Little Hercules Foundation

Little Zebra Fund

Lurie Children's Hospital

Lupus & Allied Diseases Association

Memorial Sloan Kettering Cancer Center

MORGAN Project MPS Society

Navigating Life with Genetic Mutations National Adrenal Disease Foundation

National MPS Society

National Tay-Sachs & Allied Diseases

Association

Partners for Krabbe Research

Pathways for Rare and Orphan Studies

PMD Foundation Probably Genetic Project Alive

Project 8p Foundation PTEN Foundation Rare New England Remember the Girls

Sarcoidosis of Long Island

Save BabiesThrough Screening Foundation Sickle Cell Thalassemia Patients Network

Sisterlink Collaborative, Inc

SLC6a1 Connect

Stop ALD
Taylor's Tale
Team 4 Travis
Titus Tough
The Firefly Fund

The Mount Sinai Hospital

The Myelin Project
The Jackson Project

University of California Davis Health United Leukodystrophy Foundation

University of Minnesota Masonic Children's

UNTOLD Story

Weill Cornell Medical Center

Wynne Mattefy Research Foundation Yaya Foundation for 4H Leukodystrophy