May 5, 2015

The Honorable Gus Bilirakis U.S. House of Representatives Washington, D.C. 20515

The Honorable G.K. Butterfield U.S. House of Representatives Washington, D.C. 20515

Dear Representatives Bilirakis & Butterfield:

We are writing to commend your introduction of the Orphan Product Extensions Now Accelerating Cures & Treatments, also known as the OPEN ACT HR 971. On behalf of the patients we represent, we wish to add our names in support of this important legislation.

We applaud the OPEN ACT because it will bring hundreds of safe, effective, and affordable medicines to rare disease patients within the next several years by incentivizing drug makers to "repurpose" major market drugs for the treatment of life-threatening rare diseases and pediatric cancers. Although biopharmaceutical companies are not currently repurposing major market therapies to treat rare diseases, the OPEN ACT solves this problem by making available an "Orphan Product Exclusivity Extension," which would provide an additional six months of market exclusivity to the patent life of the major market drug being repurposed so long as the sponsor company establishes that the therapy is designated to treat a rare disease and obtains a rare disease indication from the federal Food and Drug Administration (FDA) on the drug label.

With 95 percent of rare diseases having no FDA-approved cure, we are confident the OPEN ACT will result in a significant increase in the number of well-tested therapies approved by the FDA for use in treating rare disease patients. We also believe the OPEN ACT will lead to rare disease therapies priced at major market prices, fewer rare disease patients using untested and potentially ineffective drugs off–label, and a boost in investment in the biotech sector.

In closing, we thank you for championing this important piece of legislation which promises to improve the quality of life for the nearly 30 million Americans suffering from rare diseases. We are eager to work with you and your staff in advocating this important legislation and look forward to the day it is enacted.

Sincerely,

National MPS Society
With Purpose
National PKU Alliance
Taylor's Tale
RASopathies Network USA
Kids v Cancer
Let Them Be Little X2 Inc.
Info and Resources for Idiopathic Pulmonary Hemosiderosis (IPH-NET)
Noah's Hope

Mary Payton's Miracle Foundation

Hope4Bridget Foundation

Batten Disease Support & Research Association

Cure Sanfilippo Foundation

Beyond Batten Disease Foundation

Drew's Hope Scientific Research Foundation

International Pemphigus and Pemphigoid Foundation (IPPF)

Cure AHC

Autoinflammatory Alliance

MLD Foundation

Fabry Support & Information Group

Children's PKU Network

FMD Chat

National Tay-Sachs & Allied Diseases Association (NTSAD)

Little Miss Hannah Foundation

Rare Disease United Foundation

Global Genes Project

Fibromuscular Dysplasia Society of America (FMDSA)

Lymphatic Malformation Institute

Mastocytosis Society

EB Research Partnership

BRBN Alliance

Jonah's Just Begun

Abigail Alliance for Better Access to Developmental Drugs

Hannah's Hope Fund

GNE Myopathy International

The Ryan Foundation

Organic Acidemia Association

Cardio-Facio-Cutaneous International

NGLY1.org

Gwendolyn Strong Foundation

POMC Island One boy an Ocean of friends

Gene Giraffe Project

International FOP Association

Aware of Angels

CureCADASIL

GT23 FOUNDATION

Desmoid Tumor Research Foundation (DTRF)

The Association for Glycogen Storage Disease

Gene Spotlight Inc.

Amyloidosis Foundation

Heriditary Neuropathy Foundation

Relapsing Polychondritis

Klippel-Feil Syndrome Freedom

CureDuchenne

Prader-Willi Syndrome Association

EveryLife Foundation for Rare Diseases

Bert's Big Adventure

Parent Project Muscular Dystrophy

Sarcoma Foundation of America

The Nicholas Conor Institute

Luck2Tuck Foundation

Team Sanfilippo Foundation

The Rally Foundation for Childhood Cancer Research

CARES Foundation, Inc.

Help Extinguish Hunter Syndrome

Sephardic Health Organization for Referral & Education

Hunter Syndrome Research Coalition

The Kortney Rose Foundation

Saving Case & Friends

Phelan-McDermid Syndrome Foundation

The Children's Medical Research Foundation, Inc.

Cure SMA

Narcolepsy Network

Celiac Support Association

Caleb's Crusade Against Childhood Cancer

International Waldenstrom's Macroglobulinemia Foundation (IWMF)

PKD Foundation

EDSers United Foundation

Choroideremia Research Foundation, Inc.

Genetic Alliance

The Life Raft Group

The Will Luthcke Foundation

Angioma Alliance

Smashing Walnuts Foundation

Castleman Disease Collaborative Network/Castleman's Awareness & Research Effort

The GIST Cancer Awareness Foundation

The Truth 365

The Arms Wide Open Childhood Cancer Foundation

Sophia's Fund

Journey4ACure

Princesses on a Mission, Inc.

Noah's Light Foundation

Pediatric Cancer Foundation

West Virginia Kids Cancer Crusaders, Inc.

Bear Necessities Cancer Foundation

A Kids' Brain Tumor Cure

RARE Science, Inc.

ISMRD (the International Advocate for Glycoprotein Storage Diseases)

Hermansky-Pudlak Syndrome Network Inc.

Run4Rare

A-T Children's Project

The Global Foundation for Peroxisomal Disorders

The Adult Polyglucosan Body Disease Research Foundation (APBDRF)

Alexa Nawrocki Pediatric Cancer Foundation

Beckwith-Wiedemann Children's Foundation International

The Brooke Healey Foundation

Talia's Legacy Children's Cancer Foundation

The Rare Childhood Cancer Advocacy Group

Alex's Army Childhood Cancer Foundation

The Catherine Elizabeth Blair Memorial Foundation

Stillbrave Childhood Cancer Foundation

Cures Within Reach

ALL4Trey

Team Sabrina

Sofia's Hope, Inc.

ALL4Trey

Delainee's Battle

Joey's Wings Foundation

The Bozeman 3

Team Ashley Bragg

Cole vs Cancer

Dominick One in a Million

Samuel Szabo Foundation

Wilms Tumor Survivor Group

Aiden's Army

Sofia's Hope, Inc.

Mikey's Way Foundation

Team Serena

Supporting Our Cancer Kids

The Champ's Corner

Habitat for Hope

Ali's Angels Foundation

Gold Rush Cure Foundation

Sickle Cell Warriors, Inc.

The Rare Cancer Research Foundation

Carson Leslie Foundation

Amyloidosis Research Consortium

Pulmonary Fibrosis Advocates

The Coalition for Pulmonary Fibrosis

Mytonic Dystrophy Foundation

LMSarcoma Direct Research Foundation

BioPontis Alliance for Rare Diseases

Foundation for Ichthyosis & Related Skin Types, Inc.

5p- Society

The Santonio Holmes III & Long Foundation

National Fragile X Foundation

National Organization for Rare Disorders (NORD)

OsteoPETrosis Society

Curing Retinal Blindness Foundation

The MAGIC Foundation

Cure HHT

DEFY Foundation

Chase After a Cure

DC Outreach Inc.

Children's Cardiomyopathy Foundation