

August 1, 2022

The Honorable Patty Murray Chair Committee on Health, Education, Labor & Pensions United States Senate Washington, D.C. 20510

The Honorable Frank Pallone Chair Committee on Energy and Commerce United States House of Representatives Washington, D.C. 20515 The Honorable Richard Burr Ranking Member Committee on Health, Education, Labor & Pensions United States Senate Washington, D.C. 20510

The Honorable Cathy McMorris Rodgers Ranking Member Committee on Energy and Commerce United States House of Representatives Washington, D.C. 20515

Dear Chairwoman Murray, Chairman Pallone, Ranking Member Burr and Ranking Member McMorris Rodgers,

The 99 undersigned organizations, representing patients with rare diseases and other acute or chronic health conditions, urge you to resume negotiations to develop a comprehensive FDA user fee package. Our organizations are deeply concerned about the impact that a delay in passage of this legislation will have on the FDA's ability to conduct the timely review of critical products that our patients need but believe there are

other important policies that must be considered as part of the user feel reauthorization process and urge you to quickly develop robust, consensus legislation.

As you know, the Senate HELP Committee favorably reported the Food and Drug Administration Safety and Landmark Advancements (FDASLA) Act (S. 4348) out of Committee with a bipartisan 13-9 vote last month. In June, the U.S. House of Representatives considered and passed the Food and Drug Amendments of 2022 (H.R. 7667) with a bipartisan vote of 392-28. In addition to the provisions that would reauthorize FDA's user fee programs relating to prescription drugs, medical devices, biosimilars, and generic drugs, both the House and Senate bills contain additional provisions that would make necessary changes and improvements to the Federal Food, Drug, and Cosmetic Act that would ultimately benefit the patients our organizations represent. For example, both S. 4348 and H.R. 7667 would strengthen FDA's accelerated approval pathway to ensure that patients and their providers can continue to have confidence in the safety and effectiveness of drugs approved under the pathway. Furthermore, both S. 4348 and H.R. 7667 contain provisions to improve timely patient access to generic drugs and biosimilars.

Additionally, the House and the Senate bills each have distinct provisions that warrant cross-chamber consideration. For instance, H.R. 7667 includes provisions that would go a long way toward ensuring increased representation of diverse and underserved populations in the clinical trials supporting FDA-approved drugs and medical devices, but similar provisions are currently not in S. 4348. Conversely, S. 4348 includes provisions to improve FDA's oversight of the infant formula and medical food market to ensure continuous supplies of infant formula and medical foods are available that were not in the House-passed bill.

Our organizations strongly believe Congress should capitalize on the user fee reauthorization process to consider and enact the additional policies that would have little chance of passage as stand-alone bills, just as it has every five years since 1992. To pass a "clean" user fee package would be to walk away from an opportunity to make critical improvements in our nation's system for overseeing medical products. We urge you to immediately work together to blend your respective bills and continue Congress' longstanding tradition of passing strong, bipartisan FDA legislation to the benefit of all Americans. For more information, please contact Heidi Ross, Vice President of Policy and Regulatory Affairs for the National Organization for Rare Disorders, at HRoss@rarediseases.org.

Thank you for your consideration,

National Organization for Rare Disorders
Achalasia Awareness Organization
Acid Maltase Deficiency Association (AMDA)
ADNP Kids Research Foundation
Allergy Asthma & Immunology Relief
ALS Association
American Kidney Fund
Angelman Syndrome Foundation
Arthritis Foundation
Association for Creatine Deficiencies
Avery's Hope

Born a Hero, Research Foundation
California Coalition for PKU and Allied Disorders
Charcot Marie Tooth Association
Children's Tumor Foundation
Children's Cancer Cause
Cholangiocarcinoma Foundation
Chondrosarcoma CS Foundation
Choroideremia Research Foundation
Congenital Hyperinsulinism International
Cure CMD
Cure HHT

Cure VCP Disease, Inc.

Cutaneous Lymphoma Foundation

Cystic Fibrosis Research Institute (CFRI)

Dreamsickle Kids Foundation, Inc

Dup15q Alliance Epilepsy Foundation

Fabry Support & Information Group

FACES: The National Craniofacial Association Fibromuscular Dysplasia Society of America

Free ME from Lung Cancer

Friedreich's Ataxia Research Alliance

Gaucher Community Alliance Glut1 Deficiency Foundation Gorlin Syndrome Alliance GRIN2B Foundation HCU Network America

Hemophilia Federation of America

Hepatitis B Foundation

Hermansky-Pudlak Syndrome Network

Hydrocephalus Association IGA Nephropathy Foundation Immune Deficiency Foundation

International Autoimmune Encephalitis Society

International Foundation for AiArthritis

International Pemphigus Pemphigoid Foundation International Waldenstrom's Macroglobulinemia

Foundation

Juju and Friends CLN2 Warrior Foundation

Mississippi Metabolics Foundation

MLD Foundation MSA Coalition

MSUD Family Support Group Muscular Dystrophy Association National Ataxia Foundation National Brain Tumor Society National MALS Foundation

National Multiple Sclerosis Society

National PKU Alliance National PKU News

National Scleroderma Foundation

National Spasmodic Dysphonia Association

Neuromuscular Disease Foundation

No Stomach For Cancer NTM Info & Research

Organic Acidemia Association

Pheo Para Alliance

PRISMS

Pulmonary Fibrosis Foundation RASopathies Network USA

Reflex Sympathetic Dystrophy Syndrome

Association

Remember The Girls

RETpositive

Sarcoma Foundation of America

SATB2 Gene Foundation

Sickle Cell Reproductive Health Education

Directive

SLC6A1 Connect

Spina Bifida Association SSADH Association STXBP1 Foundation

Superficial Siderosis Research Alliance

SYNGAP1 Foundation

TANGO2 Research Organization

TargetCancer Foundation

Tatton Brown Rahman Syndrome Community

Team Telomere

The AKU Society of North America The Association for Frontotemporal

Degeneration (AFTD)

The Desmoid Tumor Research Foundation

The E.WE Foundation

The Global Foundation for Peroxisomal

Disorders

The LAM Foundation

The Leukemia & Lymphoma Society

The Life Raft Group
The Patient Story
The RYR-1 Foundation

Turner Syndrome Society of the United States

Vasculitis Foundation Wilson Disease Association

CC: The Honorable Charles E. Schumer, Majority Leader, U.S. Senate

The Honorable Mitch McConnell, Minority Leader, U.S. Senate

The Honorable Nancy Pelosi, Speaker of the House, U.S. House of Representatives

The Honorable Kevin McCarthy, Minority Leader, U.S. House of Representatives