April 11, 2017

The Honorable Lamar Alexander
Chairman
Senate Health, Education, Labor, and Pensions Committee
428 Senate Dirksen Office Building
Washington, DC 20510

The Honorable Patty Murray
Ranking Member
Senate Health, Education, Labor, and Pensions Committee
428 Senate Dirksen Office Building
Washington, DC 20510

The Honorable Greg Walden
Chairman
House Energy and Commerce Committee
2125 Rayburn House Office Building
Washington, D.C. 20515

The Honorable Frank Pallone
Ranking Member
House Energy and Commerce Committee
2125 Rayburn House Office Building
Washington, D.C. 20515

Dear Chairman Alexander, Ranking Member Murray, Chairman Walden, and Ranking Member Pallone:

On behalf of the Coalition for Clinical Trials Awareness, we urge you to include in the Prescription Drug User Fee Act reauthorization process a related effort to address a pressing public health challenge: lack of public awareness of the benefits of clinical trials leading to underenrollment.

Since 2013, the 40 patient, professional, and research organizations that comprise CCTA have worked to bring attention to the concerning facts about clinical trials awareness in the United States. For instance, **37% of clinical trial sites do not meet their enrollment goals.** According to the Tufts Center for the Study of Drug Development, **11% of trials fail to enroll even a single patient.**

Low participation can delay drug development and increase discovery costs, hindering patient access to life-saving treatments. Further, low enrollment of women and minorities in clinical trials makes it difficult to determine the efficacy of a medicine, and can lead to a lack of information about demographic-specific aspects of disease, outcomes, and responses to a therapy.

By working together to raise public awareness, we can improve enrollment and encourage efficient, cost-effective development of vital treatments. Toward that goal, please consider working with your colleagues to implement the following legislative measures as you consider the Prescription Drug User Fee Act reauthorization during this congressional session:
First, form an interagency task force on clinical trials awareness. The Secretary of Health and Human Services would convene a task force with representatives from the Food and Drug Administration, the National Institutes of Health, the Centers for Disease Control and Prevention, and the Department of Health and Human Services, as well as patient group representatives, health care professionals, and representatives from the pharmaceutical industry.

Second, create a roadmap based on the task force’s findings. The roadmap would describe:

- How the government, patient groups, and industry can engage in a public-private partnership to advance clinical trials awareness.
- What additional training is needed for health care professionals so that they would more appropriately inform patients about clinical trials.
- How digital technology and social media platforms can promote clinical trials awareness and participation.

Third, implement an information campaign based upon the roadmap. The Offices of the Assistant Secretary for Health and Assistant Secretary for Public Affairs would conduct an educational campaign to raise awareness of the need for and benefits of clinical trials for both patients and society at large. Demographic groups with historically low levels of clinical trials participation, including women and minorities, would be targeted by the campaign; therefore, the Offices of Women's and Minority Health would play important leadership roles in implementing this initiative. This information campaign would advance the message of clinical trials enrollment through television advertisements, social media, and outreach events for the patient and health care professional communities.

CCTA envisions what such a campaign might look like. We invite you to explore these materials at http://cctawareness.org/CTAW-2016/.

In past years, federally directed task forces, strategic plans, and information campaigns have successfully addressed important public health challenges, including drug shortages, a regulatory framework for health information technology, and the accessibility of prescription drug container information for Americans with disabilities. CCTA urges members of Congress to use the precedent set by these efforts to launch a similarly structured and equally successful initiative to improve clinical trials awareness and enrollment.

If we may be of assistance in this matter, please contact us at info@cctawareness.org or 888-507-5675.
Sincerely,

Alliance for Headache Disorders Advocacy
Alliance for Patient Access
Association of Clinical Research Organizations
Cholangiocarcinoma Foundation
Conference Forum
Fabry Support & Information Group
Gerontological Society of America
Global Colon Cancer Association
Lupus and Allied Diseases Association, Inc.
Michael J. Fox Foundation for Parkinson’s Research
Pancreatic Cancer Action Network
WomenHeart: The National Coalition for Women with Heart Disease

CC: Members, Senate Committee on Health, Education, Labor, and Pensions
     Members, House Committee on Energy and Commerce