Dear ALS Registry Participant,

The HEALEY ALS Platform Trial is enrolling people living with Amyotrophic Lateral Sclerosis (ALS) to test the efficacy of multiple investigational products. The platform trial is designed to decrease the time it takes to test new potential therapies and increase access to research for people living with ALS. One of the innovative features of the Platform Trial is that enrollment will continue as more investigational products are added.

We partner with more than 50 research centers across the USA (view site map). If you would like to enroll in this trial, please contact the research team at any local participating center (HEALEY ALS Platform Trial sites contact information). Please note that the Platform Trial involves about 6 to 8 in-person study visits over the course of 6 months, so we encourage participants to pursue enrollment at conveniently located centers.

If you have questions about the HEALEY ALS Platform Trial or would like assistance connecting with a local site, please contact the Patient Navigator via email or phone 833-425-8257 (HALT ALS).

Additionally, our weekly Q&A Webinars, held on Thursdays at 5pm ET, are a great way to stay up to date about the Platform Trial and submit questions in real time.

For more information about the Platform Trial:
- Register to attend upcoming webinars or view recordings of previous webinars:
  - Webinars
- Overview of the Platform Trial:
  - General Information: Platform Trial
  - Something New is Here: HEALEY Platform Trial
  - Frequently Asked Questions
  - What to Expect in the Trial Process
- Investigational Products in the Platform Trial:
  - Study Drugs
- Find out if you or a loved one may be eligible to participate:
  - Online Eligibility Checking Tool

Thank you,
The HEALEY ALS Platform Trial Patient Navigation Team

Please note: The National ALS Registry and ATSDR are not part of this project and have no control over it. Please directly contact the research staff listed in the attachment to this email.

If you want us to stop sending these notices, you can update the preferences on your registry account on the National ALS Registry website (www.cdc.gov/als) by logging into your account and clicking on the “Update Account” link. Then uncheck the “I Agree” box under ALS Research Notification Consent form.