Are you an ALS/MND patient or caregiver? We need your help!

Take a survey to help guide drug development and services for amyotrophic lateral sclerosis (ALS)/motor neurone disease (MND)

Please access this important survey at:

In an effort to include your perspective to help guide the development of new drugs and services for ALS, we are conducting a survey of patients with ALS and of caregivers. This initiative, titled IMPACT-ALS (Investigating and Measuring Patient and Caregiver Trends about ALS), is a research collaboration between industry, European ALS doctors and ALS charities.
The survey is intended to identify the burden of ALS and concerns during the course of the disease, preferences for treatment perceptions of patients and caregivers. Results from this survey are planned to be published in peer-reviewed journals and other channels. The results of the survey will be accessible to all participants.

The survey will be available from 13/10/2020 to 20/11/2020

✓ The survey should take approximately 30–45 minutes to complete
✓ Responses are completely anonymous

If you have any questions or difficulty while completing the survey, please contact: ALS_Survey_Europe@apothecom.com

The information that you provide in this survey will:

- Be anonymised. You will not be identifiable
- Be used solely for the purpose of research studies relating to ALS treatments and experiences of ALS patients and caregivers
- Improve future experiences for other ALS patients and caregivers
- Be shared with companies who have helped to fund the study
- Not be shared with companies if it is personal information, and you will not be contacted by anybody from industry about the information you have provided if you participate in this study