

CLINICAL DEVELOPMENT SOLUTIONS TO IMPROVE AMYOTROPHIC LATERAL SCLEROSIS RESEARCH

Given the rapidly progressing nature of amyotrophic lateral sclerosis (ALS) and the lack of effective treatments or cures, the need for new treatment options for this rare disease is urgent. Signant Health can support the drug development process by applying our SmartSignals clinical research solutions as well as scientific expertise to help researchers improve trial outcomes.

IMPROVED PATIENT EXPERIENCE

Reduce burdens for your ALS study participants while simultaneously generating better-quality PRO data. SmartSignals eCOA helps study teams implement PRO assessments electronically so patients can self-report without the burden of travel. Plus, it supports caregiver-assisted completion with built-in attributability features.

OPTIMIZED ENDPOINT RELIABILITY

Ensure investigative staff administer and score ClinRO assessments accurately by leveraging our clinician-developed Rater Training system as well as our PureSignal Analytics solution. We'll train raters and then monitor data in real-time, providing opportunities to proactively correct errors or other quality concerns before they impact the reliability of your endpoint data.

PATIENT-FOCUSED REMOTE SOLUTIONS

Make study participation more convenient for patients suffering from symptoms of ALS. With eConsent, patients can take their time reviewing study requirements at home before making the effort to visit the clinic. Once enrolled, your team can leverage our secure, compliant Telemedicine platform to connect with patients and caregivers between visits or for appointments not requiring site-based assessments.

EXPERT CLINICAL ADVISORY

With decades of clinical science experience, our ALS research leaders are prepared to serve as an extension of your study team. Rely on our in-house neurology experts to help optimize ALS protocols, implement clinically-appropriate instruments, and oversee data quality from launch through regulatory submission.

AGILE RTSM

Simplify randomization for multi-arm trial designs and minimize product waste and overages with Signant's agile RTSM solution. We'll help you launch faster, manage mid-study changes with ease, and support your trial operations no matter how or where you conduct them.

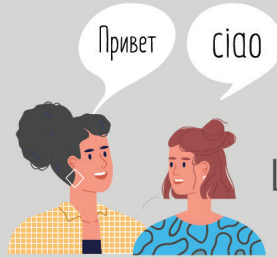
DIGITALIZE THE PROCESS

Each solution and service within our SmartSignals evidence generation platform can be applied to a study independently. However, when combined, they create an intuitive and powerful digital ecosystem for creating and managing complex global studies. Plus, every study is supported by a dedicated team of clinical science and operations experts.

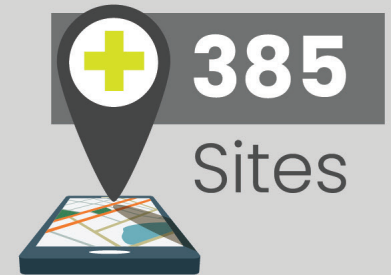
At Signant, our focus is helping you develop and deliver treatments or therapies faster in order to improve the quality of life for ALS patients everywhere.

SIGNANT'S ALS CLINICAL TRIAL EXPERIENCE

Phases



15
Languages



385
Sites



10
Countries



2,100
Patients

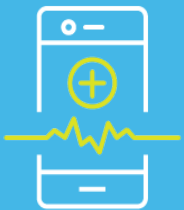


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Studies

SMARTSIGNALS SOLUTIONS

The SmartSignals solutions can be used individually or integrated together for a seamless, end-to-end digital experience.

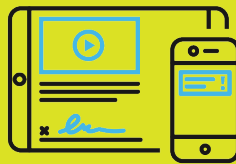
eCOA



Scientific Consulting & Advisory



eConsent



Rater Training & PureSignal Analytics



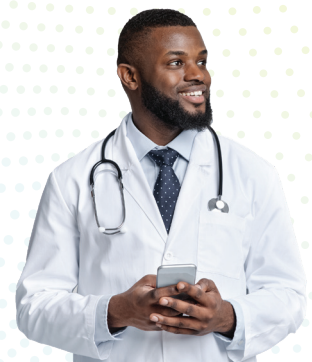
RTSM



DISCUSS YOUR NEXT STUDY WITH US

Our global team of therapeutic area experts advise on all areas of the clinical development process, including:

- Clinical science and medicine
- Data analysis
- Regulatory
- Operations and trial administration
- Global logistics



MEET THE EXPERTS