SIGNANT SMARTSIGNALS® UNIFIED PLATFORM



CLINICAL TRIAL TECHNOLOGY SOLUTIONS AND SERVICES TAILORED FOR PHASE I CLINICAL TRIALS



Phase I clinical trials provide vital early information on the safety and tolerability of new medicines, and help to inform early dosage and regimen considerations. Studies are typically small, and start and complete rapidly, with Phase I units operating several studies concurrently. Technology solutions offer important operational efficiencies, but need to support rapid implementation and end-to-end processes. Signant's technology solutions, including our Unified Platform EDC with add-on modules for eConsent, Randomization, eCOA, and TeleVisits, provide the perfect partner for Phase I research.

Signant's Unified Technology Solution Addresses Phase I Trial Requirements:



Comprehensive Technology Suite

Our Signant SmartSignals[®] Unified Platform contains all the essential technology components for Phase I research, fully integrated. Leverage our comprehensive EDC and include add-on modules for eConsent, Randomization, eCOA and TeleVisits as needed.



Rapid Set Up

Rapid full platform solution set up. EDC in typically 4-6 weeks or less, using our intuitive designer that requires no-software coding. Our extensive library of re-usable forms and study templates speeds study solution assembly.



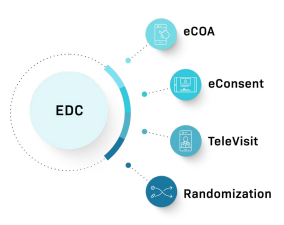
Simplicity for the Site

Site users love our truly unified platform. From a single sign on, sites can access all studies and all applications. Applications are integrated out-of-the-box, and site users enjoy seamless, converged workflow between applications. Simplify data collection with direct data capture using our mobile-optimized eSource interface.

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Reduced Training Burden

Our Unified Platform solutions are easy to use and intuitive. No need to learn multiple vendor solutions – all solutions are part of the same, single platform.



Powered by Our Unified eClinical Platform

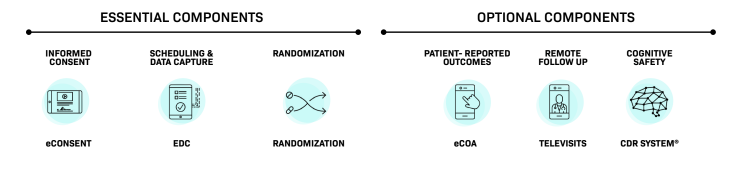
Rapid set-up, comprehensive EDC, with add-on modules for eCOA, eConsent, TeleVisits and Randomization. Our modular approach puts you in control, with a robust EDC foundation and optional add-ons tailored to your study needs.

Built on a single database, there is no need to provide back-end integrations between platform solutions.

Phase I units can build their own study solutions using self-service, or Signant Health's experienced teams can provide full-service including study solution design and implementation.

Why Partner with Signant Health?

Our unified solution suite provides a comprehensive end-to-end solution for Phase I research.



INFORMED CONSENT

Using eConsent:

- Easily upload pdf ICF documents and include multimedia resources
- Provide participant access to study information and ICF ahead of / during site visit
- Improve transparency and quality of consenting documentation and eliminate inspection findings

SCHEDULING AND DATA CAPTURE

Using EDC:

- Leverage direct data capture, and reduce SDV, through modern responsive design that can be used on any mobile device
- Access study level and cross-study scheduling and activity calendars
- Enable remote SDV to reduce on-site CRA visits

RANDOMIZATION

Using Randomization:

- Include randomization within EDC workflow one click randomization from the same common, intuitive interface
- · Access rapid and transparent emergency unblinding

PATIENT-REPORTED OUTCOMES

Using eCOA:

 Easily collect integrated ePRO data when needed by the study – such as pharmacodynamic measures, formulation acceptability and taste, and reactogenicity

REMOTE FOLLOW UP

Using TeleVisits:

 Enable engaging follow up assessments using video meetings between site and participant

COGNITIVE SAFETY

Using Signant SmartSignals[®] CDR System[®]:

 Assess treatment-related changes in attention and processing speed

WHO IS SIGNANT HEALTH?



Signant Health is the evidence generation company. We are focused on leveraging software, deep therapeutic and scientific knowledge, and operational expertise to consistently capture, aggregate, and reveal quality evidence for clinical studies across traditional, virtual, and hybrid trial models. For more than 25 years, over 600 sponsors and CROs of all sizes – including all Top 20 pharma – have trusted Signant solutions for remote and site-based eCOA, EDC, eConsent, RTSM, supply chain management, and data quality analytics. Learn more at www.signanthealth.com.