

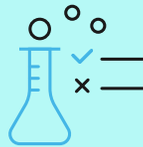
RELIABLE EVIDENCE FOR MEDICAL DEVICE PRODUCT APPROVALS

The pathway to approval for new diagnostic and therapeutic products, or combination drug/device products, isn't always straightforward. Signant blends proven evidence generation and trial optimization solutions with clinical and operational expertise to help you meet your protocol and product goals.

From endpoint selection to regulatory submission, Signant's solutions and resources help device studies succeed with:



Proven evidence generation and clinical trial optimization technologies



Industry-leading clinical and eCOA science expertise



Global scale & operational capacity

WHY PARTNER WITH SIGNANT FOR MEDICAL DEVICE PRODUCT TRIALS?

- 01** We invest time to understand your study goals and longer-term product objectives
- 02** A comprehensive array of eClinical technologies, backed by scientific and operational expertise, are at your disposal from one partner
- 03** Our infrastructure, resources, as well as country and site intelligence support global trials at any scale
- 04** Test your products across a variety of therapeutic areas and indications in consultation with our clinical science and medicine team
- 05** We stay up-to-date on regulatory changes such as increased emphasis on the use of patient-reported outcomes in medical device trials
- 06** As eCOA experts, we ensure you choose study-appropriate instruments as well as advise on measurement and data capture strategies

STREAMLINING THE PATHWAY TO APPROVAL WITH SIGNANT SMARTSIGNALS SOLUTIONS

DISCOVER HOW OUR END-TO-END SOLUTION SUITE OPTIMIZES PRODUCT RESEARCH PARTICIPATION AND OPERATIONS.



COA DATA CAPTURE

Capture accurate, complete COA data with our eCOA solutions



ENDPOINT SELECTION & PROTOCOL DESIGN

Leverage our clinical science and medicine expertise throughout a study's lifecycle



REMOTE STUDY VISITS

Optimize remote study operations with our comprehensive telemedicine platform



PATIENT ENGAGEMENT

Keep patients on-track and engaged



INVENTORY MANAGEMENT

Get products where they are needed with agile, quick-deploy RTSM



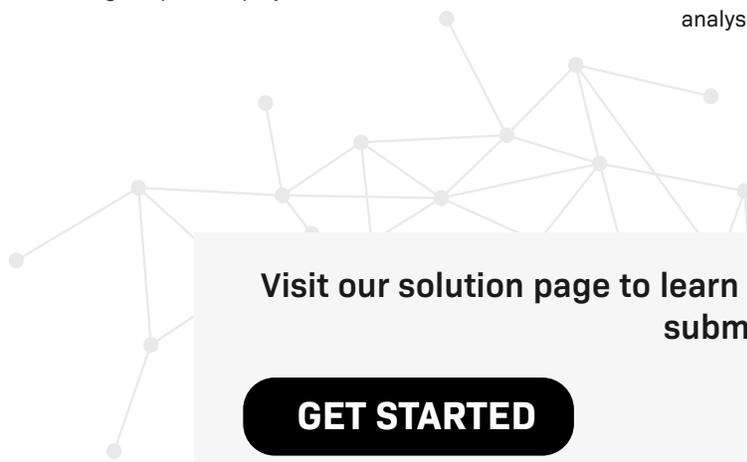
CLINICAL DATA MANAGEMENT

Capture, aggregate, and standardize clinical data for reviews and analyses



PARTICIPANT CONSENT

Simplify the process of obtaining informed consent and re-consent remotely or at site



Visit our solution page to learn more and submit an RFP.

GET STARTED

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 20 years, over 400 sponsors and CROs of all sizes - including all Top 20 pharma - have trusted Signant solutions for remote and site-based eCOA, eConsent, RTSM, supply chain management, and data quality analytics. Learn more at www.signanthealth.com.