

When the World is Counting on You, Count on Signant to Help Deliver Your Vaccine

In the high-stakes world of vaccine development, every moment and data point counts. Vaccine studies can be large, multinational endeavors with sensitive timeframes, extended participant follow up periods, multiphase designs, and critical data management needs.



Signant's science-led solutions help sponsors and CROs overcome the core challenges of vaccine trials by streamlining high-quality data collection, enhancing participant engagement, and simplifying trial operations—ultimately accelerating your path to market while maintaining the highest standards of quality and compliance.



eCOA

- Capture critical safety data with our validated reactogenicity diary, with adult and pediatric versions
- Drive accurate efficacy assessment with long-term disease surveillance diaries
- Provisioned and BYOD options
- Drive high compliance with alarms, reminders, and site notifications
- Meet delivery milestones for seasonal disease studies



RTSM

- Scale to handle high numbers of patients randomized rapidly
- Flexible medication vial usage rules (single or multiple participants)
- Seamless adaptation for multi-phase designs
- Full cold-chain capabilities including quarantine management
- Transparent split shipment management for frozen and non-frozen supplies



eConsent

Simplify your consent process with our flexible options tailored to your study design



Telemedicine

Unburden long-term participant follow-up for disease incidence assessment without requiring site visits



Patient Engagement

Guide participants through the study, provide clinic reminders, and maintain engagement in long-term surveillance periods

SIGNANT SMARTSIGNALS® Vaccine Trial Solutions



Proven Under Pressure

When speed and scale were critical, we delivered. During the race to develop a COVID-19 vaccine, we helped launch a 44,000-participant global trial across 150 sites in seven countries in just five weeks, supporting 150 sites with our solutions and round-the-clock operational support. The result? Our sponsor achieved both Emergency Use Authorization and full regulatory approval in record time while maintaining the highest standards of data quality.

REIMAGINE THE PATH TO PROOF WITH SIGNANT HEALTH.

Driving High Completion Compliance

Example: RSV Vaccine Trial

38,000+

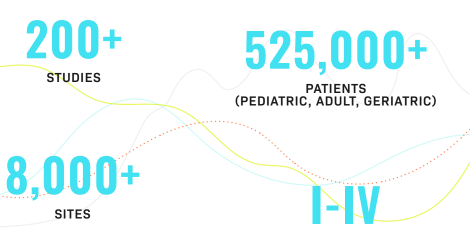
60+ YEARS

DEMOGRAPHICS

86.6%

Signant's Vaccine Study Experience

PHASES

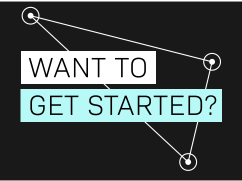


30+

OUNTRIES

50+

LANGUAGES



DISCUSS PROTOCOL

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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.