

Veramed

Confidence in your clinical trials

Clinical trials are high stakes, and high stakes demand high standards.

You need a biometrics CRO that cares about your trials as much as you do. One that brings together the right minds, culture and systems to deliver biometrics solutions you never have to second guess.

From end-to-end support to rapid-response study rescue, we tailor our solutions and work as an extension of your team to drive your studies and empower your decisions.

“Veramed is the number 1 CRO that we have ever worked with. The quality and communication we have received has been excellent and we definitely want to work with Veramed on all our studies moving forward.”

— Biotech client

Services

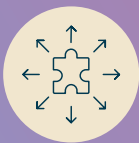
Data Management
Biostatistics
Programming
Standards, Innovation & AI
Evidence & Value Generation

Solutions

Strategic & Technical Consulting
Project-Based Solutions
Functional Service Provider



Dedicated,
collaborative
support



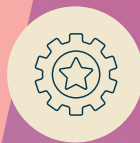
Scalable,
tailored
solutions



Accelerated
timelines



Trusted, strategic
leadership and
governance



Innovative
processes and
deep expertise



Regulatory
confidence

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Success stories



Detective work in data: CDISC conversion excellence

Across six months, Veramed led a data forensics mission, converting legacy data dating back to 2011 into CDISC-compliant formats, as well as identifying and reconstructing missing data. Veramed's meticulous work and rigorous validation processes delivered high-quality, regulatory-ready outputs under complex and evolving circumstances.

From rescue to FDA approval: Reanalysis and resubmission of an ophthalmology trial

A client with a novel ophthalmological product for a condition with very limited treatment options reached out to Veramed after failings with the original CRO. As a result of Veramed's statistical re-analysis of the data and new submission, this novel treatment received FDA approval. Efficiently getting the analysis back on track accelerated the time to market for an advanced treatment that could have a life-changing impact for affected patients.

Advancing an orphan drug: Agility, collaboration and rare disease expertise

A biotech client needed a CRO that could offer better communication, prioritization and leadership than their current vendor. Veramed deployed a team with a proven track record in rare disease studies, along with a structured transition plan. As a smaller, more agile organization, Veramed's team could rapidly adapt to client requirements and respond efficiently to questions that were raised, forming the collaborative extension of the client's team that was desired.

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Veramed
Biometrics Built Better