

COMPANY PROFILE



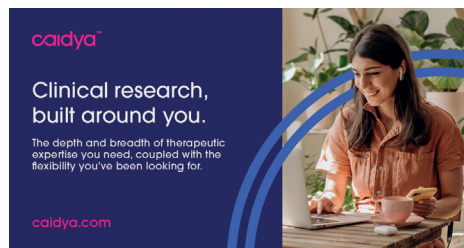
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History and Mission

As leaders in research, we are entrusted with important work. So naturally, we think it is important to provide an environment where our employees know their ideas, opinions and contributions all matter. We value transparency and integrity and support a performance-oriented culture where everyone is encouraged to collaborate. We want our employees to know that working here is more than a just job – it's an opportunity to change lives and contribute to the future of medicine.

Highlights

Caidya is a multi-therapeutic clinical research organization (CRO) serving innovators worldwide. Focused on delivery excellence and an elevated customer experience, Caidya offers a wide range of clinical services and vast therapeutic expertise, supporting its partners from pre-IND strategy, through clinical development to submission and post-marketing surveillance. Caidya leverages industry-leading and proprietary clinical technology to ensure trial transparency and data-driven decision-making. Formed in 2021 following the merger of leading CROs, dMed and Clinipace, Caidya has nearly 1,800 employees in 30+ countries throughout the world.



Caidya

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BVMA member since 2004
Audits passed in 2004, 2006, 2009,
2012, 2015, 2018 and 2022



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Services

Clinical Development Phase I-IV

- Biostatistics and statistical programming
- Clinical data management
- Clinical operations
- Clinical technology
- Investigator recruitment and site selection
- Medical monitoring
- Medical writing
- Project management
- Protocol feasibility
- Quality assurance
- Regulatory affairs
- Safety/pharmacovigilance

Therapeutic Areas

- Gastroenterology, nephrology and urology, oncology, rare disease, and women's health
- Cardiology, CNS, ENT, immunology, infectious diseases and vaccines, nutrition, orthopaedic, and respiratory

Regulatory Affairs

- Product development
- Portfolio management
- Regulatory affairs
- Manufacturing and quality support
- Medical and regulatory writing
- Regulatory publishing/submissions