

## COMPANY PROFILE



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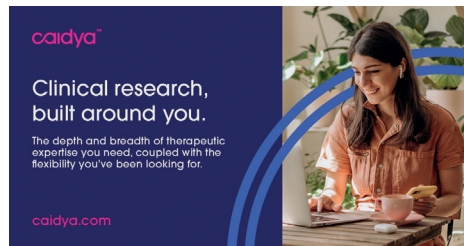
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## History and Mission

**Caidya's** mission is to help bring life-changing therapies to our global community through liberated clinical research. We believe in creating innovations that improve the overall development process and help bring quality treatments to patients more quickly and effectively. Caidya began in 2003 as a software company under the brand name Clinipace, and through a series of global acquisitions and mergers became a full-service CRO.

## 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 1700 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



[www.caidya.com](http://www.caidya.com)

## 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
- Safety/pharmacovigilance

### Therapeutic Areas

- Oncology, Rare Disease, Immunology, Neurology, Infectious Diseases; Cardiovascular, Dermatology, Healthy Volunteers, Hematology, Ophthalmology, Endocrinology, Gastroenterology, Women's Health

### Regulatory Affairs

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



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