

EMPOWER YOUR PORTFOLIO

Your CRO Service Partner

Why together

Close cooperation between Sponsors and CROs is a well-established routine. Yet, even with a shared understanding of how study planning and execution should work, typical hurdles and challenges remain. From a CRO perspective, it can sometimes feel as if Sponsors expect teams to be everywhere at once, especially when timelines are tight and milestones must be met. This is where pressure builds and delivering quality in time gets challenging.

Camovis is the right partner to stay on track. We step in to fill operational gaps and strengthen teams from the very beginning to the final close-out of a clinical trial, always on demand and precisely where support is needed most.

We enable. We speed up.

- **Since 2014**, we have been supporting sponsors, CROs, and trial sites in the execution of clinical trials according to the latest regulatory requirements.
- **Over 350** carefully selected Camovis research professionals operate throughout Germany, Switzerland, Austria and beyond.
- **We are dedicated to delivering.** Anytime, anywhere, on-site and remote. Camovis is your partner for high-performance, patient-centered clinical trial services.

Our Commitment — We plug in

“Camovis complements existing operational structures and accelerates study execution. We are committed to be the perfect partner for CROs and their teams to deliver clinical trials smooth and successfully.”

Carolyn Kurth | CEO | Camovis

- Close stakeholder collaboration
- Deep integration in site operations
- Focus on timing and quality
- Acceleration of operational workflows
- Bridging HR & resource gaps
- Engaged, agile, and patient-centric

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Camovis Services

We are where clinical trials take place: on-site at the trial centers. At the same time, we support teams in study organization, empower staff through training, and provide consulting when processes or recruitment face obstacles. In this way, we combine operational services, training, management, and consulting to ensure that studies are not only planned but also successfully executed.

Site Coordination:	Operational management of study workflows.
Patient Recruitment:	Targeted identification, pre-screening, and follow-up.
Clinical Operations:	Conduct of study visits, medical examinations, and sample logistics.
Data Handling:	Data transfer, eCRF entry, query management, and archival data.
DCT Services:	Home visits, remote support, virtual visits, and patient-centered care.
Project Training:	Practical, study-focused training for site teams.
Courses:	Webinars and e-learning for study coordinators and study nurses.
Consulting:	Site performance enhancement, audit & inspection readiness, and coaching.

Let`s get to work

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Feel free to contact us — we're here and happy to help! :)

