



AXONCRO

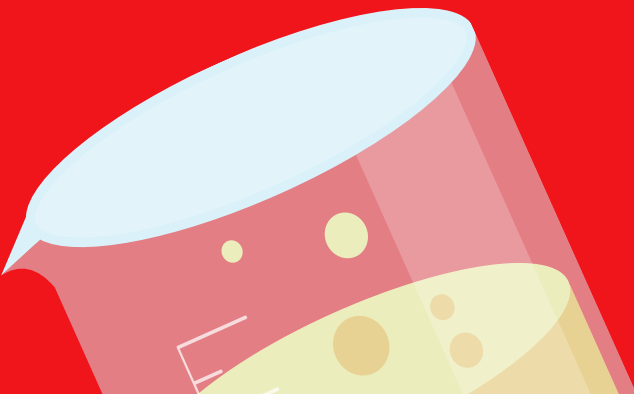
PRESENTATION
FOR NEW
COLLABORATIONS



Resolute Supporter At Clinical Trials



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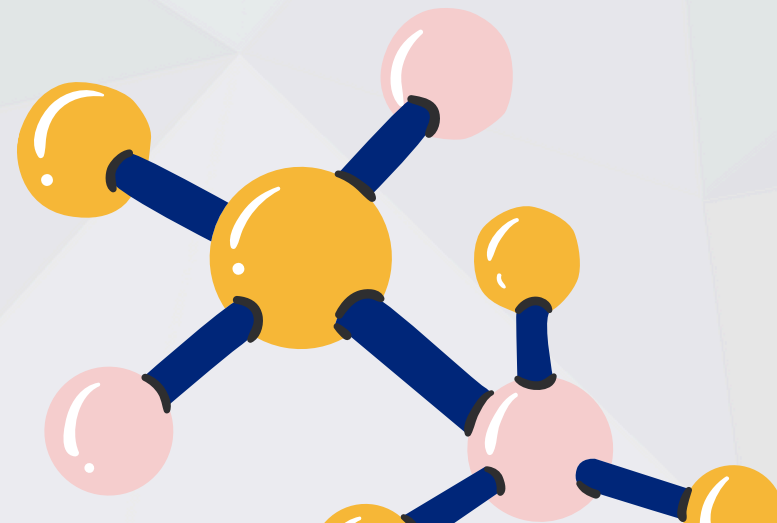


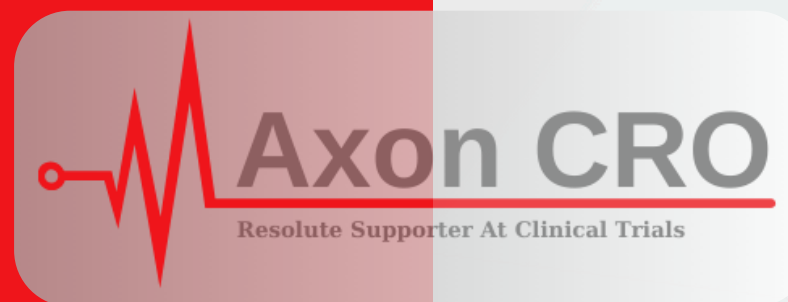


Who We Are

Axoncro is a reliable CRO that bridges Turkey's clinical research potential with global standards. We deliver end-to-end solutions – from protocol design to final reporting.

With nationwide operational capability, we proudly conduct clinical studies across 5 provinces and 8 active centers, leveraging Turkey's vast investigator and volunteer potential.





Mission & Vision

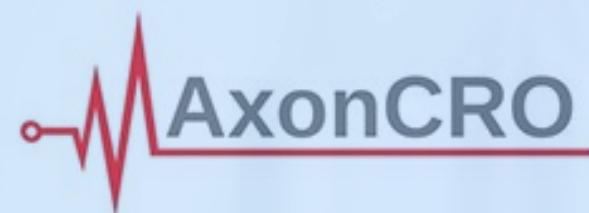
Our Mission

To empower pharmaceutical companies in their clinical research journey by offering reliable, ethical, and scientifically robust solutions – ensuring that every project is successfully executed in compliance with international standards.

Our Vision

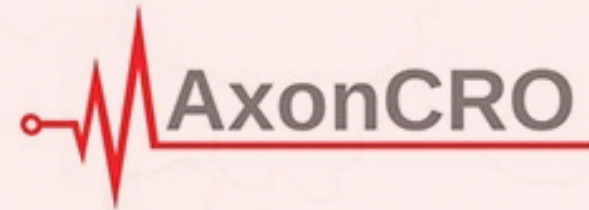
To be recognized as a leading and trusted CRO in Turkey, and the preferred strategic partner for global and local sponsors.





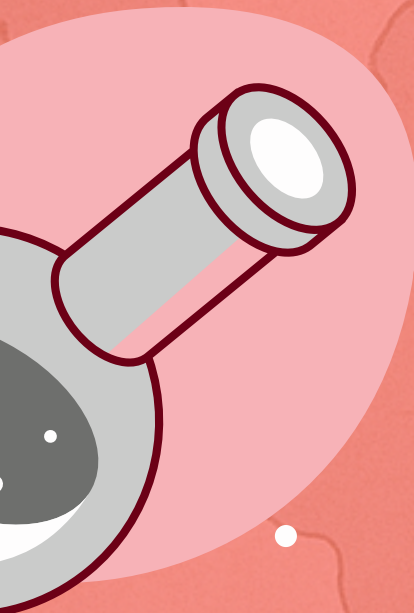
Why Choose Axoncro?

-  Nationwide Reach – 5 provinces, 8 active centers
-  Strong Investigator Network – Experienced researchers & motivated volunteers
-  Ethics & Reliability – Patient safety is our top priority
-  End-to-End Services – All processes under one roof
-  Global Standards – Full compliance with ICH-GCP & national regulations
-  Academic Collaboration – Training the next generation of professionals



Why Conduct Clinical Trials in Türkiye?

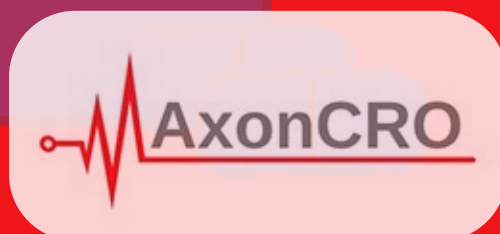
- - ◆ Diverse Patient Population
 - Access to a large, demographically diverse patient pool
 - Enables faster recruitment and more representative study outcomes
 - ◆ Experienced Research Teams
 - Skilled investigators, coordinators, and site staff with international trial expertise
 - Ensures scientific rigor and high-quality data collection
 - ◆ Dedicated Clinical Trial Units
 - Specialized departments in hospitals and healthcare facilities
 - Organized, structured, and seamless trial execution
 - ◆ Efficient Regulatory Pathways
 - Transparent and well-structured approval processes
 - Faster study start-up compared to many other regions
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Our Services



- Clinical Trial Management – Protocol development, site selection, patient recruitment
- Site Management (SMO) – Coordination of study centers, staff support
- Monitoring & Data Management – On-site monitoring, electronic data capture and reporting
- Regulatory & Ethical Consulting – Guidance through national and international requirements
- Medical Writing & Translation – Clinical documentation, publications, regulatory texts
- Personnel & Training – CRA, coordinators, and experienced site staff





Our Services

Your Trusted Partner in Clinical Research

At the heart of our work is a commitment to advancing science and improving patient care. We specialize in the design and management of clinical drug trials, observational studies, medical device research, and early access programs for humanitarian use—always ensuring excellence and compliance at every step.

◆ End-to-End Expertise

From feasibility planning to local authority approvals, site audits, and final study close-out, we deliver seamless clinical trial management tailored to your needs.

◆ Strong Investigator Network

With our extensive and experienced investigator network, we provide accurate feasibility assessments and build reliable foundations for successful studies.

◆ Flawless Site Operations

Our dedicated site coordinators ensure all processes are carried out smoothly and in full compliance with ICH-GCP principles, local regulations, and study protocols.

◆ Comprehensive Support

We go beyond coordination by offering data entry services, clinical research nurse support, and full operational management, ensuring your studies run efficiently and without disruption.

✨ With deep sector experience and a passion for quality, we position ourselves as your strategic partner in clinical research success.





🎓 Bridging Academia & Industry

🎓 Academic Collaboration & Sector Development

At Axoncro, we not only manage clinical research projects but also invest in the future of the sector.

Through collaboration with Kocaeli Health and Technology University, we contribute to clinical research education, equipping the next generation of professionals with both academic knowledge and practical skills.



Who We Partner With

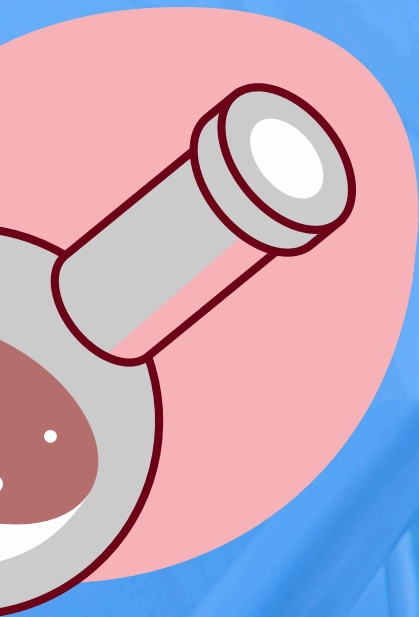
Axoncro is committed to serving:

- Global and local pharmaceutical companies
- Biotech firms advancing innovative therapies
- Medical device manufacturers
- Academic institutions seeking CRO support





Contact Us



AXONCRO Headquarters
Yahya Kaptan Mah. Şehit Ergün Köncü Sk. No: 9/A Kat 1
İzmit / Kocaeli – Turkey

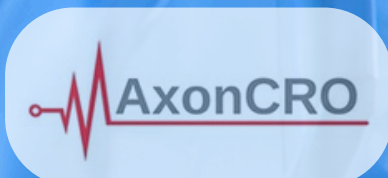
✉ mehmetacar@axoncro.com

✉ esraacar@axoncro.com

☎ +90 505 731 80 25

☎ +90 531 620 59 90

www.axoncro.com



CLOSING NOTE

WITH NATIONWIDE CAPABILITY, STRONG SCIENTIFIC AND ETHICAL STANDARDS, AND TURKEY'S CLINICAL RESEARCH POTENTIAL – AXONCRO IS YOUR UNWAVERING ALLY IN ADVANCING INNOVATIVE TREATMENTS.

 “YOUR DEDICATED PARTNER IN CLINICAL RESEARCH.”

THANK YOU FOR YOUR ATTENTION

