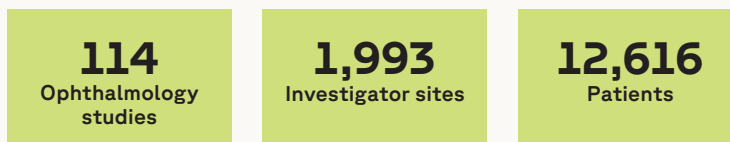


Drive your ophthalmic device development with an experienced partner.

Overcoming development challenges and bringing a medical device to market relies on strategic insights and dedicated experience. Learn how our team at Fortrea is uniquely equipped to provide agility at scale and help advance life-changing ophthalmic devices—from the clinic to approval.

Apply our ophthalmic clinical experience

With nearly four decades of experience in ophthalmology since 1985, we've earned the trust of sponsors around the world. In the last five years, we've been selected as the CRO of choice to support:



Access a global CRO with local, scalable resources

With Fortrea as your dedicated partner, you can leverage our strong familiarity with the scientific, investigator and patient resources across Asia Pacific, Europe, North America, Latin America and Middle East/Africa.

Whether you need support with your protocol, trial design or post-market study, we can offer a range of solutions tailored to your product's specific needs.

- Clinical study, strategy and operations
- Clinical events committee management
- Clinical compliance/auditing
- Protocol development
- Patient recruitment and retention
- Site recruitment and qualification
- Site management and site monitoring
- Database development and management
- Data management
- Biostatistics consulting
- Data Monitoring Committee (DMC)/ Data Safety Monitoring Board (DSMB) Management
- Safety services
- Statistical design and analysis
- Vendor management

Look ahead at regulatory and commercialization challenges

We're here to navigate regulations while meeting your clinical and commercialization objectives. Throughout your ophthalmic device development journey, we offer our expertise in global regulatory strategies—including reimbursement strategies and commercialization—to help you maximize the potential of your device.

Sharpen your focus on quality

Our team has invested in key certification standards such as ISO 13485 to help minimize audit complexity and maximize quality compatibility throughout your program. And, as with any high-performance team, we strive to form an interactive, transparent relationship to achieve greater results together.

Our strategic vision for your ophthalmic device

At Fortrea, we know how to form a strong clinical partnership aligned to your requirements. With our dedicated medical device team, we're focused on delivering high-quality and agile services that align to your protocol and flexibly scale to your needs. Together, we believe the exceptional is possible.



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