

K233421 RESCAN 700Mar 8, 2024
150 days to decisionK233421 · Product code: **OBO** · Ophthalmic
Source: <https://www.510kdatabase.net/k233421/>**SUBMISSION DETAILS**

| | |
|-----------------------|-------------------------------------|
| Decision | Substantially Equivalent (Cleared) |
| Submission type | Traditional |
| Device classification | Tomography, Optical Coherence (OBO) |
| Date received | Oct 10, 2023 |
| Decision date | Mar 8, 2024 |
| Days to decision | 150 days |
| Third-party review | No |
| Combination product | No |
| PCCP authorized | No |
| Summary / Statement | Summary |

APPLICANT

| | |
|----------------|---|
| Company | Carl Zeiss Meditec, AG |
| Location | Dublin, CA, US |
| Contact | Katrin Faber |
| Website | http://www.zeiss.com/meditec-ag/en_de/home.html |
| 510(k) history | 45 submissions · 44 cleared · 2004-2025 |

Carl Zeiss Meditec, AG is a global medical device manufacturer specializing in innovative solutions for ophthalmology and microsurgery. The company operates with a manufacturing facility in Dublin, US, and delivers diagnostic and surgical instruments to healthcare professionals worldwide. The company has received FDA 510(k) clearances from total submissions since 2004. Ophthalmic devices represent the dominant category, accounting for 71% of submissions. The latest clearance in 2025 reflects continued regulatory activity and product innovation in this specialized field. C...

REGULATORY CONSULTANT

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|-----------------|---------------------------------|
| Consulting firm | Carl Zeiss Meditec, Inc. |
| Contact | Aditya Rao |

Regulatory consulting firm that managed this 510(k) submission on behalf of the applicant. Source: [FDA accessdata.fda.gov](https://accessdata.fda.gov)
