

K082016 GDX VCC WITH ECC SOFTWAREAug 10, 2009
391 days to decisionK082016 · Product code: **MYC** · Ophthalmic
Source: <https://www.510kdatabase.net/k082016/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Ophthalmoscope, Laser, Scanning (MYC)
Date received	Jul 15, 2008
Decision date	Aug 10, 2009
Days to decision	391 days
Third-party review	No
Summary / Statement	Summary

APPLICANT

Company	Carl Zeiss Meditec, Inc.
Location	San Diego, CA, US
Contact	JUDITH A BRIMACOMBE
Website	https://www.zeiss.com/meditec
510(k) history	29 submissions · 29 cleared · 1993-2025

Carl Zeiss Meditec, Inc. is a global medical device manufacturer specializing in innovative solutions for ophthalmology and microsurgery. The company operates with a manufacturing facility in San Diego, California, and is part of the ZEISS Group, a leader in optical and optoelectronic technologies since 1846. The company maintains a strong FDA 510(k) regulatory record with FDA 510(k) clearances from total submissions. Ophthalmic devices represent the dominant focus, accounting for approximately 86% of submissions. Carl Zeiss Meditec has been active in FDA clearances since...
