

K895538 ANGIOSCOPE W/THE OPTISCOPE-2 INTEGRATED ANGIOSCOPYApr 26, 1990
225 days to decisionK895538 · Product code: LYK · Cardiovascular
Source: <https://www.510kdatabase.net/k895538/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Angioscope (LYK)
Date received	Sep 13, 1989
Decision date	Apr 26, 1990
Days to decision	225 days
Third-party review	No

APPLICANT

Company	Trimedyne, Inc.
Location	Walker, MI, US
Contact	MERRITT M GIRGIS
Website	http://www.trimedyne.com/
510(k) history	58 submissions · 58 cleared · 1981-2005

Trimedyne, Inc. is a manufacturer of Holmium:YAG lasers and surgical peripherals. The company specializes in laser-based surgical solutions for minimally invasive procedures across multiple specialties including urology, orthopedics, spine surgery, and general surgery. Trimedyne has received FDA 510(k) clearances from total submissions since its first clearance in 1981. The company's regulatory focus centers on General & Plastic Surgery devices, which represent 83% of its submission history. The latest clearance on record dates to 2005, reflecting the company's historical...

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