

**K840142 YAG LASER 8000 - LOWER GI APPLICAT**May 18, 1984  
133 days to decisionK840142 · Product code: **LNK** · Gastroenterology & UrologySource: <https://www.510kdatabase.net/k840142/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Laser For Gastro-urology Use (LNK)
Date received	Jan 6, 1984
Decision date	May 18, 1984
Days to decision	133 days
Third-party review	No

**APPLICANT**

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Company	<b>Cooper Lasersonics, Inc.</b>
Location	Orangeburg, NY, US
Website	<a href="https://www.cooperhealthcare.com">https://www.cooperhealthcare.com</a>
510(k) history	54 submissions · 52 cleared · 1982-1988

Cooper Lasersonics, Inc. is a medical device manufacturer based in Orangeburg, US, specializing in laser surgical systems. The company has received FDA 510(k) clearances from total submissions since its first clearance in 1982. Cooper Lasersonics focused primarily on CO2 and Nd:YAG laser systems for surgical applications across multiple specialties, including general and plastic surgery, otolaryngology, gastroenterology, and neurosurgery. The company's regulatory activity concluded in 1988, making this a historical record of its contributions to surgical laser technology....

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