

K170121 Lumenis Family of Holmium Surgical Lasers and Delivery Devices and Accessories

May 22, 2017
129 days to decisionK170121 · Product code: **GEX** · General & Plastic SurgerySource: <https://www.510kdatabase.net/k170121/>

SUBMISSION DETAILS

Decision	Substantially Equivalent (Cleared)
Submission type	Special
Device classification	Powered Laser Surgical Instrument (GEX)
Date received	Jan 13, 2017
Decision date	May 22, 2017
Days to decision	129 days
Third-party review	No
Summary / Statement	Summary

APPLICANT

Company	Lumenis, Ltd.
Location	Santa Clara, CA, US
Contact	Yehudit Kraizer
Website	http://www.lumenis.com/
510(k) history	28 submissions · 27 cleared · 2003-2021

Lumenis, Ltd. is a global leader in energy-based medical device solutions for aesthetic and vision care. The company develops and commercializes innovative laser, intense pulsed light (IPL), and radiofrequency technologies with a manufacturing facility in Santa Clara, California. Lumenis has received FDA 510(k) clearances from total submissions since 2003. The company specializes in General & Plastic Surgery devices, which represent 93% of its regulatory submissions. Its cleared portfolio includes ablative and fractional CO₂ laser systems, IPL platforms, and surgical lase...

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Device record: <https://www.510kdatabase.net/k170121/>. Data sourced from FDA 510(k) public records (accessdata.fda.gov).

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