

K850175 MENTOR DBL-COIL INDWELL. SILICONE URETERAL STENT SMar 7, 1985
49 days to decisionK850175 · Product code: **FAD** · Gastroenterology & Urology
Source: <https://www.510kdatabase.net/k850175/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Stent, Ureteral (FAD)
Date received	Jan 17, 1985
Decision date	Mar 7, 1985
Days to decision	49 days
Third-party review	No

APPLICANT

Company	Mentor Corp.
Location	Mchenry, IL, US
Contact	GREGORY L JOHNSON
510(k) history	61 submissions · 61 cleared · 1977-2013

Mentor Corp. is a surgical aesthetics and medical device company based in McHenry, US. Now part of Johnson & Johnson MedTech, the brand supplies products to plastic surgeons and specialists worldwide. Mentor has received FDA 510(k) clearances from total submissions since its first clearance in 1977. The company's regulatory record spans General & Plastic Surgery, Gastroenterology & Urology, Obstetrics & Gynecology, and Radiology device categories. The latest clearance was recorded in 2013, reflecting the company's historical significance in surgical device innovation. Men...
