

K780901 AUTOMATED ELECTROPHORESIS SYSTEMJun 28, 1978
27 days to decisionK780901 · Product code: **JQT** · Chemistry
Source: <https://www.510kdatabase.net/k780901/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Densitometer/scanner (integrating, Reflectance, Tlc, Radiochromat.) Clinica (JQT)
Date received	Jun 1, 1978
Decision date	Jun 28, 1978
Days to decision	27 days
Third-party review	No

APPLICANT

Company	Olympus Corp.
Location	Mchenry, IL, US
Website	https://www.olympus-global.com
510(k) history	142 submissions · 140 cleared · 1978-1995

Olympus Corp. is a Japanese optics and imaging manufacturer founded in 1919. The company operates from McHenry, US, and holds approximately 70 percent of the global endoscope market. Olympus has received FDA 510(k) clearances from total submissions between 1978 and 1995. The company's cleared devices span gastroenterology, urology, obstetrics, gynecology, and chemistry specialties. This regulatory record reflects the company's historical focus on endoscopic surgical instruments and related technologies. Notable cleared device categories include resection electrodes, fiber...