

K820111 SIEMENS SIRESKOP 4Feb 18, 1982
34 days to decisionK820111 · Product code: **KXJ** · Radiology
Source: <https://www.510kdatabase.net/k820111/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Table, Radiologic (KXJ)
Date received	Jan 15, 1982
Decision date	Feb 18, 1982
Days to decision	34 days
Third-party review	No

APPLICANT

Company	Siemens Corp.
Location	Mchenry, IL, US
Website	http://www.siemens.it/
510(k) history	66 submissions · 66 cleared · 1978-2010

Siemens Corp. is a global technology company headquartered in McHenry, US. The company develops medical imaging and diagnostic equipment for healthcare providers worldwide. Siemens has received FDA 510(k) clearances from total submissions. The company's regulatory focus centers on Radiology devices, which represent the dominant category of its cleared portfolio. FDA 510(k) clearances span from 1978 to 2010, establishing a significant historical record in medical device regulation. Recent cleared devices include advanced imaging systems such as CT scanners, MR systems, X-r...
