

**K960382 KODAK DIGITAL SCIENCE MEDICAL IMAGE  
MANAGER**Apr 17, 1996  
83 days to decisionK960382 · Product code: **LMD** · Radiology  
Source: <https://www.510kdatabase.net/k960382/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	System, Digital Image Communications, Radiological (LMD)
Date received	Jan 25, 1996
Decision date	Apr 17, 1996
Days to decision	83 days
Third-party review	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Eastman Kodak Company</b>
Location	Mchenry, IL, US
Contact	NANCY BUTCHER
Website	<a href="http://www.kodak.com">http://www.kodak.com</a>
510(k) history	238 submissions · 238 cleared · 1977-2006

Eastman Kodak Company is a diversified imaging and materials manufacturer headquartered in McHenry, US. The company has a long history in advanced materials, chemicals, and imaging technologies. Eastman Kodak maintains a significant regulatory history in medical imaging devices. The company received FDA 510(k) clearances from total submissions, with clearances spanning from 1977 to 2006. The company's cleared devices focused primarily on radiology and medical imaging systems, including digital radiography systems, picture archiving and communication systems (PACS), and re...

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Device record: <https://www.510kdatabase.net/k960382/> Data sourced from FDA 510(k) public records ([accessdata.fda.gov](http://accessdata.fda.gov)).

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