

**K822774 9110 SYRINGE**Nov 3, 1982  
50 days to decisionK822774 · Product code: **FMF** · General Hospital  
Source: <https://www.510kdatabase.net/k822774/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Syringe, Piston (FMF)
Date received	Sep 14, 1982
Decision date	Nov 3, 1982
Days to decision	50 days
Third-party review	No

**APPLICANT**

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Company	<b>Cardiac Pacemakers, Inc.</b>
Location	Mchenry, IL, US
510(k) history	76 submissions · 76 cleared · 1977-2010

Cardiac Pacemakers, Inc. (CPI), doing business as Guidant Cardiac Rhythm Management, manufactured implantable cardiac rhythm management devices. Now part of Boston Scientific, the company is based in Saint Paul, Minnesota, with historical operations in McHenry, US. The company received FDA 510(k) clearances from total submissions between 1977 and 2010. Cardiovascular devices dominated the regulatory portfolio at 83% of submissions. This historical record reflects the company's core focus on cardiac rhythm management and related interventional technologies. CPI developed t...

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