

**K933014 LIFESTREAM CENTRIFUGEL PUMP INTERFACE  
#210PI**Sep 28, 1994  
464 days to decisionK933014 · Product code: **KFM** · Cardiovascular  
Source: <https://www.510kdatabase.net/k933014/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Pump, Blood, Cardiopulmonary Bypass, Non-roller Type (KFM)
Date received	Jun 21, 1993
Decision date	Sep 28, 1994
Days to decision	464 days
Third-party review	No
Summary / Statement	Statement

**APPLICANT**

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Company	<b>St. Jude Medical, Inc.</b>
Location	Salt Lake City, UT, US
Contact	JOSEPH MAGLIOZZI
Website	<a href="http://www.sjm.com/">http://www.sjm.com/</a>
510(k) history	23 submissions · 22 cleared · 1989-2018

St. Jude Medical, Inc. was a global medical device company headquartered in Little Canada, Minnesota. The company operated more than 20 principal facilities worldwide and sold products in over 100 countries. St. Jude Medical received FDA 510(k) clearances from total submissions between 1989 and 2018. The company specialized exclusively in Cardiovascular devices, establishing a focused portfolio in cardiac monitoring, catheter systems, and related interventional technologies. Founded in 1976 and publicly listed in 1977, St. Jude Medical achieved Fortune 500 status annually...

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