

**K082304 ROLL-X GUIDEWIRE**Aug 27, 2008  
14 days to decisionK082304 · Product code: **DQX** · Cardiovascular  
Source: <https://www.510kdatabase.net/k082304/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Wire, Guide, Catheter (DQX)
Date received	Aug 13, 2008
Decision date	Aug 27, 2008
Days to decision	14 days
Third-party review	Yes
Summary / Statement	Summary

**APPLICANT**

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Company	<b>St Jude Medical</b>
Location	Minnetonka, MN, US
Contact	JEFF STURM
Website	<a href="http://www.sjm.com/">http://www.sjm.com/</a>
510(k) history	105 submissions · 105 cleared · 2000-2018

St Jude Medical 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 2000 and 2018. The company's regulatory focus centered on Cardiovascular devices, which represented 91% of all submissions. Notable cleared products include cardiac mapping systems, pacing catheters, and mobile cardiac applications. Now part of Abbott Laboratories following its acquisition in Janua...

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