

**K132008 6981M LEAD EXTENDER KIT, 6984M LEAD EXTENDER KIT, 6986M LEAD EXTENDER KIT, 5866-24M LEAD ADAPTOR KIT, 5866-38M LEAD ADAP**Jul 30, 2013  
29 days to decisionK132008 · Product code: **DTD** · Cardiovascular  
Source: <https://www.510kdatabase.net/k132008/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Special
Device classification	Pacemaker Lead Adaptor (DTD)
Date received	Jul 1, 2013
Decision date	Jul 30, 2013
Days to decision	29 days
Third-party review	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Medtronic, Inc.</b>
Location	Mounds View, MN, US
Contact	MICHELE MACHACEK
Website	<a href="https://www.medtronic.com">https://www.medtronic.com</a>
510(k) history	209 submissions · 208 cleared · 1981-2026

Medtronic, Inc. is a global medical device manufacturer headquartered in Mounds View, United States. The company develops and markets a broad range of medical devices across multiple therapeutic areas. Medtronic maintains a strong FDA 510(k) regulatory track record with FDA 510(k) cleared devices from total submissions since 1981. The company specializes primarily in Cardiovascular devices, which represent 82% of its submission portfolio. Recent clearances include coronary perfusion cannulae, intracoronary shunts, venous cannulae, guidewires, deflectable catheter systems,...