

**K250199 VitalFlow Console**May 20, 2025  
117 days to decisionK250199 · Product code: **QNR** · Cardiovascular  
Source: <https://www.510kdatabase.net/k250199/>**SUBMISSION DETAILS**

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|                       |  |
|-----------------------|--|
| Decision              | Substantially Equivalent (Cleared)                   |
| Submission type       | Traditional  |
| Device classification | Blood Pump For Ecmo, Long-term (> 6 Hours) Use (QNR) |
| Date received         | Jan 23, 2025   |
| Decision date         | May 20, 2025   |
| Days to decision      | 117 days   |
| Third-party review    | No   |
| Combination product   | No   |
| PCCP authorized       | No   |
| Summary / Statement   | Summary  |

**APPLICANT**

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|----------------|---|
| Company        | <b>Medtronic, Inc.</b>  |
| Location       | Mounds View, MN, US   |
| Contact        | Elizabeth Rose  |
| 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,...

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510k Database - [www.510kdatabase.net](http://www.510kdatabase.net)

Device record: <https://www.510kdatabase.net/k250199/> Data sourced from FDA 510(k) public records ([accessdata.fda.gov](http://accessdata.fda.gov)).

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