

**K151715 T3 Software**Oct 29, 2015  
126 days to decisionK151715 · Product code: **MWI** · Cardiovascular  
Source: <https://www.510kdatabase.net/k151715/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Monitor, Physiological, Patient (without Arrhythmia Detection Or Alarms) (MWI)
Date received	Jun 25, 2015
Decision date	Oct 29, 2015
Days to decision	126 days
Third-party review	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Etiometry, Inc.</b>
Location	Boston, MA, US
Contact	DIMITAR BARONOV
Website	<a href="https://www.etiometry.com">https://www.etiometry.com</a>
510(k) history	11 submissions · 11 cleared · 2015-2026

Etiometry, Inc. is a clinical decision-support software company founded in 2010. Based in Boston, the company develops AI-driven clinical intelligence platforms for intensive care settings. The platform aggregates patient data, provides risk analytics, automates clinical pathways, and supports quality improvement initiatives in high-acuity care environments. Etiometry has received FDA 510(k) clearances from total submissions since 2015. The company specializes exclusively in Cardiovascular devices and software solutions. The latest clearance in 2026 demonstrates continued...