

**K143369 MRidium 3860+ MRI infusion Pump/ Monitoring System**Dec 15, 2016  
751 days to decisionK143369 · Product code: **FRN** · General Hospital  
Source: <https://www.510kdatabase.net/k143369/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Pump, Infusion (FRN)
Date received	Nov 25, 2014
Decision date	Dec 15, 2016
Days to decision	751 days
Third-party review	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Iradimed Corporation</b>
Location	Winter Park, FL, US
Contact	FRANCIS X CASEY
Website	<a href="http://www.iradimed.com/">http://www.iradimed.com/</a>
510(k) history	7 submissions · 7 cleared · 2005-2025

Iradimed Corporation specializes in MRI-safe medical devices for patient care in magnetic resonance imaging environments. The company develops infusion pumps, patient monitoring systems, and ferrous metal detection solutions designed for use during MRI procedures. Iradimed operates with a manufacturing facility in Winter Park, Florida. The company maintains a strong FDA 510(k) regulatory record with FDA 510(k) clearances from total submissions since its first clearance in 2005. All submissions have resulted in clearances, with no denials. The latest clearance in 2025 demo...

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

Device record: <https://www.510kdatabase.net/k143369/> Data sourced from FDA 510(k) public records ([accessdata.fda.gov](http://accessdata.fda.gov)).

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