

**K163331 Discovery MR750 3.0T**Mar 17, 2017  
109 days to decisionK163331 · Product code: **LNH** · Radiology  
Source: <https://www.510kdatabase.net/k163331/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	System, Nuclear Magnetic Resonance Imaging (LNH)
Date received	Nov 28, 2016
Decision date	Mar 17, 2017
Days to decision	109 days
Third-party review	No
Summary / Statement	Summary
Other names	Discovery MR750w 3.0T;Discovery MR450 1.5T; Discovery MR450w 1.5T; SIGNA Architect and SIGNA Artist

**APPLICANT**

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Company	<b>Ge Medical Systems, LLC</b>
Location	Waukesha, WI, US
Contact	Mary A. Mayka
Website	<a href="https://www.gehealthcare.com">https://www.gehealthcare.com</a>
510(k) history	104 submissions · 104 cleared · 2003-2026

GE Medical Systems, LLC is a medical device manufacturer based in Waukesha, US. The company specializes in Radiology devices and solutions. GE Medical Systems has received FDA 510(k) clearances from total submissions. The company's regulatory focus is entirely on Radiology devices, with a clearance history spanning from 2003 to 2026. The latest clearance in 2026 demonstrates active regulatory engagement within the past two years. Recent cleared devices include advanced imaging systems such as the Photonova Spectra series, SIGNA™ product line, AIR Recon DL, Revolution Vibe...

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