

K260724 MAGENTIQ-COLO (ME-APDS)Apr 3, 2026
29 days to decisionK260724 · Product code: **QNP** · Gastroenterology & Urology
Source: <https://www.510kdatabase.net/k260724/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Special
Device classification	Gastrointestinal Lesion Software Detection System (QNP)
Date received	Mar 5, 2026
Decision date	Apr 3, 2026
Days to decision	29 days
Third-party review	No
Combination product	No
PCCP authorized	No
Summary / Statement	Summary

APPLICANT

Company	Magentiq Eye, Ltd.
Location	Haifa, IL
Contact	Dror Zur
510(k) history	4 submissions · 4 cleared · 2023-2026

REGULATORY CONSULTANT

Consulting firm	Hogan Lovells US LLP
Contact	John J. Smith

Regulatory consulting firm that managed this 510(k) submission on behalf of the applicant. Source: FDA accessdata.fda.gov

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