

**K243807 Single Use Retrieval Basket V FG-
V421PR/V422PR/V431P/V432P**Mar 18, 2025
97 days to decisionK243807 · Product code: LQR · Gastroenterology & Urology
Source: <https://www.510kdatabase.net/k243807/>**SUBMISSION DETAILS**

| | |
|-----------------------|------------------------------------|
| Decision | Substantially Equivalent (Cleared) |
| Submission type | Traditional |
| Device classification | Dislodger, Stone, Biliary (LQR) |
| Date received | Dec 11, 2024 |
| Decision date | Mar 18, 2025 |
| Days to decision | 97 days |
| Third-party review | No |
| Combination product | No |
| PCCP authorized | No |
| Summary / Statement | Summary |

APPLICANT

| | |
|----------------|---|
| Company | Olympus Medical Systems Corp. |
| Location | Hachiochi-Shi, JP |
| Contact | Seiko Yunoki |
| Website | https://www.olympus-global.com |
| 510(k) history | 101 submissions · 101 cleared · 2012-2026 |

Olympus Medical Systems Corp. is a global medical device manufacturer headquartered in Hachiochi-Shi, Japan. The company specializes in endoscopic imaging systems and therapeutic devices for minimally invasive procedures. Olympus has received FDA 510(k) clearances from total submissions since 2012. The company's regulatory portfolio is dominated by Gastroenterology & Urology devices, including endoscopes, hemostatic forceps, biopsy instruments, and sphincterotomes. The latest clearance in 2026 reflects continued active development and market engagement. Recent cleared dev...

REGULATORY CONSULTANT

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|-----------------|--|
| Consulting firm | Olympus Corporation of the Americas |
| Contact | Roshana Ahmed |

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

510k Database - www.510kdatabase.net

Device record: <https://www.510kdatabase.net/k243807/> Data sourced from FDA 510(k) public records (accessdata.fda.gov).

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