

K253448 Sonosite MT Ultrasound System

Nov 3, 2025
32 days to decision

K253448 · Product code: IYN · Radiology
Source: <https://www.510kdatabase.net/k253448/>

SUBMISSION DETAILS

| | |
|-----------------------|---|
| Decision | Substantially Equivalent (Cleared) |
| Submission type | Traditional |
| Device classification | System, Imaging, Pulsed Doppler, Ultrasonic (IYN) |
| Date received | Oct 2, 2025 |
| Decision date | Nov 3, 2025 |
| Days to decision | 32 days |
| Third-party review | Yes |
| Combination product | No |
| PCCP authorized | No |
| Summary / Statement | Summary |

APPLICANT

| | |
|----------------|---|
| Company | FUJIFILM Sonosite, Inc. |
| Location | Bothell, WA, US |
| Contact | Anoush Frankian |
| Website | https://www.sonosite.com |
| 510(k) history | 25 submissions · 25 cleared · 2013-2026 |

FUJIFILM Sonosite, Inc. is a portable ultrasound manufacturer based in Bothell, US. The company specializes in point-of-care ultrasound systems for clinical imaging. FUJIFILM Sonosite has received FDA 510(k) clearances from total submissions since 2013. The company's portfolio focuses exclusively on Radiology devices. The latest clearance was in 2026, demonstrating continued regulatory activity and product innovation. The company's cleared devices include portable ultrasound systems designed for diverse clinical settings. Products span multiple system lines, each configur...

REGULATORY CONSULTANT

| | |
|-----------------|--|
| Consulting firm | Regulatory Technology Services, LLC |
| Contact | Prithul Bom |

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