

K240720 TiLink-P SI Joint Fusion SystemApr 11, 2024
27 days to decisionK240720 · Product code: **OUR** · Orthopedic
Source: <https://www.510kdatabase.net/k240720/>**SUBMISSION DETAILS**

| | |
|-----------------------|------------------------------------|
| Decision | Substantially Equivalent (Cleared) |
| Submission type | Special |
| Device classification | Sacroiliac Joint Fixation (OUR) |
| Date received | Mar 15, 2024 |
| Decision date | Apr 11, 2024 |
| Days to decision | 27 days |
| Third-party review | No |
| Combination product | No |
| PCCP authorized | No |
| Summary / Statement | Summary |

APPLICANT

| | |
|----------------|---|
| Company | SurGenTec, LLC |
| Location | Boca Raton, FL, US |
| Contact | Guilherme Pires |
| Website | https://www.surgentec.com |
| 510(k) history | 23 submissions · 23 cleared · 2017-2026 |

SurGenTec, LLC is a medical device manufacturer specializing in orthopedic surgical solutions. The company operates with a manufacturing facility in Boca Raton, US. SurGenTec has received FDA 510(k) clearances from total submissions since its first clearance in 2017. Orthopedic devices represent 78% of the company's regulatory portfolio. The company remains actively engaged in FDA 510(k) submissions, with its most recent clearance in 2026. SurGenTec's product portfolio includes fusion systems, graft delivery instruments, bone void fillers, and specialized surgical navigat...