

K250155 Xpert KneeApr 23, 2025
92 days to decisionK250155 · Product code: **HRS** · Orthopedic
Source: <https://www.510kdatabase.net/k250155/>**SUBMISSION DETAILS**

| | |
|-----------------------|------------------------------------|
| Decision | Substantially Equivalent (Cleared) |
| Submission type | Traditional |
| Device classification | Plate, Fixation, Bone (HRS) |
| Date received | Jan 21, 2025 |
| Decision date | Apr 23, 2025 |
| Days to decision | 92 days |
| Third-party review | No |
| Combination product | No |
| PCCP authorized | No |
| Summary / Statement | Summary |

APPLICANT

| | |
|----------------|---|
| Company | Newclip Technics |
| Location | Haute-Goulaine, FR |
| Contact | Gaëlle Gourbière |
| Website | https://www.newcliptechnics.com |
| 510(k) history | 31 submissions · 31 cleared · 2006-2026 |

Newclip Technics is a French manufacturer of orthopedic osteosynthesis solutions based in Haute-Goulaine, near Nantes. The company designs, manufactures, and markets surgical instruments and implants for hospitals and clinics worldwide. The company has received FDA 510(k) clearances from total submissions since its first clearance in 2006. All submissions focus on orthopedic devices. The latest clearance in 2026 confirms the company remains actively engaged in regulatory submissions and product innovation. Recent cleared devices include patient-matched instrumentation sys...

REGULATORY CONSULTANT

| | |
|-----------------|----------------------------------|
| Consulting firm | The Ortomedix Group, Inc. |
| Contact | J.D. Webb |

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