

**K231069 PERLA® TL Posterior Thoraco-lumbar Fixation System**Oct 25, 2023  
194 days to decisionK231069 · Product code: **NKB** · Orthopedic  
Source: <https://www.510kdatabase.net/k231069/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Thoracolumbosacral Pedicle Screw System (NKB)
Date received	Apr 14, 2023
Decision date	Oct 25, 2023
Days to decision	194 days
Third-party review	No
Combination product	No
PCCP authorized	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Spineart SA</b>
Location	Plan-Les-Ouates, CH
Contact	Franck Pennesi
Website	<a href="https://www.spineart.com">https://www.spineart.com</a>
510(k) history	11 submissions · 11 cleared · 2019-2026

Spineart SA is a Swiss-based orthopedic medical device company founded in 2005. Headquartered in Geneva with a manufacturing facility in Plan-Les-Ouates, Switzerland, the company specializes in innovative spine surgery solutions. Their portfolio includes motion preservation technologies, posterior fixation systems, interbody fusion devices, and enabling surgical technologies. Spineart has received FDA 510(k) clearances from total submissions since 2019. The company focuses exclusively on orthopedic devices, with a strong emphasis on minimally invasive spine surgery instru...