

**K251335 Tera Lumbar Interbody Fusion System (Various PNs)**Jun 2, 2025  
33 days to decisionK251335 · Product code: **MAX** · Orthopedic  
Source: <https://www.510kdatabase.net/k251335/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Intervertebral Fusion Device With Bone Graft, Lumbar (MAX)
Date received	Apr 30, 2025
Decision date	Jun 2, 2025
Days to decision	33 days
Third-party review	No
Combination product	No
PCCP authorized	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Acuity Surgical Devices, LLC</b>
Location	Irving, TX, US
Contact	Chuck Forton
Website	<a href="https://acuitysurgical.com">https://acuitysurgical.com</a>
510(k) history	8 submissions · 8 cleared · 2021-2025

Acuity Surgical Devices, LLC develops spinal implant solutions for surgeons and patients. Based in Irving, Texas, the company delivers complete systems for lumbar, cervical, and biologic spine surgery since 2013. The company has received FDA 510(k) clearances from total submissions, all in Orthopedic devices. Clearances span 2021 to 2025, demonstrating sustained regulatory activity and market presence in spinal implant technology. Acuity's product portfolio includes stand-alone anterior lumbar fusion systems, modular cervical fixation platforms, posterior lumbar interbody...