

**K183705 IdentiTi Porous Ti Interbody System**Mar 1, 2019  
60 days to decisionK183705 · Product code: **MAX** · Orthopedic  
Source: <https://www.510kdatabase.net/k183705/>**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	Dec 31, 2018
Decision date	Mar 1, 2019
Days to decision	60 days
Third-party review	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Alphatec Spine, Inc.</b>
Location	Carlsbad, CA, US
Contact	Cynthia Adams
Website	<a href="https://www.alphatecspine.com">https://www.alphatecspine.com</a>
510(k) history	93 submissions · 93 cleared · 2005-2026

Alphatec Spine, Inc. is a spine surgery medical device company based in Carlsbad, California. The company develops and markets surgical solutions for spinal fusion and fixation procedures. Alphatec Spine maintains a strong FDA 510(k) regulatory record with FDA 510(k) clearances from total submissions. The company specializes in Orthopedic devices, which represent 91% of its submission portfolio. Clearances span from 2005 to 2026, demonstrating sustained regulatory activity and recent market engagement. Recent cleared devices include robotic navigation systems, interbody s...

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