

K191906 MiRus 3D Printed Lumbar Interbody Fusion Systems consisting of the Callisto 3D Printed PLIF, HYPERION 3D Printed TLIF, CALYPSO 3D Printed LLIF, and ANTARES 3D Printed ALIF

May 18, 2020
307 days to decision

K191906 · Product code: **MAX** · Orthopedic
Source: <https://www.510kdatabase.net/k191906/>

SUBMISSION DETAILS

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Intervertebral Fusion Device With Bone Graft, Lumbar (MAX)
Date received	Jul 16, 2019
Decision date	May 18, 2020
Days to decision	307 days
Third-party review	No
Combination product	No
PCCP authorized	No
Summary / Statement	Summary

APPLICANT

Company	MiRus, LLC
Location	Marietta, GA, US
Contact	Jordan Bauman
Website	https://www.mirusmed.com
510(k) history	24 submissions · 24 cleared · 2018-2026

MiRus, LLC is a medical device company based in Marietta, Georgia. The company develops innovative orthopedic implants and surgical solutions. MiRus has received FDA 510(k) clearances from total submissions since its first clearance in 2018. The company specializes exclusively in orthopedic devices, with a focus on spinal fusion systems, interbody fusion devices, and osteotomy solutions. Recent clearances include posterior cervical fusion systems, lumbar plating systems, and expandable wedge osteotomy devices. The company remains actively engaged in FDA submissions, with ...

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Device record: <https://www.510kdatabase.net/k191906/> Data sourced from FDA 510(k) public records (accessdata.fda.gov).

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