

K211186 Erchonia XLR8Oct 22, 2021
185 days to decisionK211186 · Product code: **NHN** · Physical Medicine
Source: <https://www.510kdatabase.net/k211186/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Powered Light Based Laser Non-thermal Instrument With Non-heating Effect For Adjunctive Use In Pain Therapy (NHN)
Date received	Apr 20, 2021
Decision date	Oct 22, 2021
Days to decision	185 days
Third-party review	No
Combination product	No
PCCP authorized	No
Summary / Statement	Summary

APPLICANT

Company	Erchonia Corporation
Location	Fountain Inn, SC, US
Contact	Steven Shanks
Website	https://www.erchonia.com
510(k) history	26 submissions · 26 cleared · 2010-2026

Erchonia Corporation is a medical device manufacturer based in Fountain Inn, US, specializing in low-level laser technology for therapeutic applications. The company has received FDA 510(k) clearances from total submissions, with a regulatory track record spanning 2010 to 2026. Erchonia's cleared devices focus primarily on General & Plastic Surgery and Physical Medicine applications, including laser systems for pain management, fat loss, and tissue healing. The company remains actively engaged in FDA regulatory submissions, with its most recent clearance in 2026. Erchonia...

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Device record: <https://www.510kdatabase.net/k211186/> Data sourced from FDA 510(k) public records (accessdata.fda.gov).

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