

K251012 TRUDERMAL Pro (ZLD-390)Aug 14, 2025
134 days to decisionK251012 · Product code: **OHS** · General & Plastic Surgery
Source: <https://www.510kdatabase.net/k251012/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Light Based Over The Counter Wrinkle Reduction (OHS)
Date received	Apr 2, 2025
Decision date	Aug 14, 2025
Days to decision	134 days
Third-party review	No
Combination product	No
PCCP authorized	No
Summary / Statement	Summary

APPLICANT

Company	Shenzhen Kaiyan Medical Equipment Co., Ltd.
Location	Shenzhen, CN
Contact	Dijkstra Alain
Website	https://www.kaiyanmedical.com
510(k) history	21 submissions · 21 cleared · 2023-2026

Shenzhen Kaiyan Medical Equipment Co., Ltd. is a medical device manufacturer based in Shenzhen, China. The company specializes in light therapy devices for clinical and aesthetic applications. Kaiyan Medical has received FDA 510(k) clearances from total submissions since 2023. All cleared devices fall within the General & Plastic Surgery category. The company maintains active regulatory status, with its most recent clearance in 2026. The company's cleared device portfolio includes LED light therapy masks, hair growth helmets, and advanced light-based aesthetic treatment s...