

K243848 Cala kIQ

Jan 16, 2025
31 days to decision

K243848 · Product code: **QBC** · Neurology
Source: <https://www.510kdatabase.net/k243848/>

SUBMISSION DETAILS

Decision	Substantially Equivalent (Cleared)
Submission type	Special
Device classification	External Upper Limb Tremor Stimulator (QBC)
Date received	Dec 16, 2024
Decision date	Jan 16, 2025
Days to decision	31 days
Third-party review	No
Combination product	No
PCCP authorized	No
Summary / Statement	Summary

APPLICANT

Company	Cala Health, Inc.
Location	Burlingame, CA, US
Contact	Alexander Kent
Website	https://www.calahealth.com
510(k) history	7 submissions · 6 cleared · 2018-2026

Cala Health, Inc. develops wearable neuromodulation devices for tremor management. The company operates with a manufacturing facility in Burlingame, California. Cala’s flagship technology delivers transcutaneous afferent patterned stimulation (TAPS) therapy to reduce hand tremor in patients with essential tremor and Parkinson’s disease. The company has received FDA 510(k) clearances from total submissions since its first clearance in 2018. All submissions focus on Neurology devices. The most recent clearance was in 2026, confirming active regulatory engagement and ongoing...

REGULATORY CONSULTANT

Consulting firm	Gardner Law PLLC
Contact	Amanda Johnston

Regulatory consulting firm that managed this 510(k) submission on behalf of the applicant. Source: [FDA accessdata.fda.gov](https://accessdata.fda.gov)
