

K241390 NeuroMatchNov 26, 2024
195 days to decisionK241390 · Product code: **OMB** · Neurology
Source: <https://www.510kdatabase.net/k241390/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Automatic Event Detection Software For Full-montage Electroencephalograph (OMB)
Date received	May 15, 2024
Decision date	Nov 26, 2024
Days to decision	195 days
Third-party review	No
Combination product	No
PCCP authorized	No
Summary / Statement	Summary

APPLICANT

Company	LVIS Corporation
Location	Palo Alto, CA, US
Contact	Sweta Srivastava
510(k) history	3 submissions · 3 cleared · 2023-2025

REGULATORY CONSULTANT

Consulting firm	Mcra, LLC
Contact	Devjani Saha

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

510k Database - www.510kdatabase.netDevice record: <https://www.510kdatabase.net/k241390/> Data sourced from FDA 510(k) public records (accessdata.fda.gov).

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