

K970491 DUPEL II BUFFERED IONTOPHORESIS ELECTRODESMay 30, 1997
109 days to decisionK970491 · Product code: **EGJ** · Physical Medicine
Source: <https://www.510kdatabase.net/k970491/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Device, Iontophoresis, Other Uses (EGJ)
Date received	Feb 10, 1997
Decision date	May 30, 1997
Days to decision	109 days
Third-party review	No
Summary / Statement	Summary

APPLICANT

Company	Empi
Location	Walker, MI, US
Contact	STACY MATTSON
Website	http://www.empik.com
510(k) history	60 submissions · 56 cleared · 1977-2010

Empi is a historical medical device manufacturer based in Walker, US. The company specialized in electrotherapy and neurostimulation devices. Empi received FDA 510(k) clearances from total submissions between 1977 and 2010. The company's primary focus was physical medicine and neurology devices, including transcutaneous nerve stimulators, iontophoresis systems, and electrotherapy equipment. This regulatory record spans over three decades of device development and market clearance. Empi is now a historical record only, with no FDA 510(k) clearances issued in more than 13 y...
