

K831787 PATIENT MONITOR #78353ASep 29, 1983
118 days to decisionK831787 · Product code: **DRT** · Cardiovascular
Source: <https://www.510kdatabase.net/k831787/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Monitor, Cardiac (incl. Cardiotachometer & Rate Alarm) (DRT)
Date received	Jun 3, 1983
Decision date	Sep 29, 1983
Days to decision	118 days
Third-party review	No

APPLICANT

Company	Hewlett-Packard Co.
Location	Mchenry, IL, US
Website	https://www.hp.com
510(k) history	230 submissions · 229 cleared · 1976-2000

Hewlett-Packard Co. is a technology company headquartered in McHenry, US. The company historically developed medical devices alongside its core computing and printing business. Hewlett-Packard received FDA 510(k) clearances from total submissions, with clearances spanning 1976 to 2000. The company specialized in cardiovascular devices, including defibrillators, telemetry systems, and clinical information systems. Additional cleared devices covered gastroenterology, urology, and radiology applications. This regulatory record reflects the company's historical involvement in...

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