

**K770756 ECHOCARDIOGRAPHY SYS., MODEL 3860**May 9, 1977  
14 days to decisionK770756 · Product code: **DXK** · CardiovascularSource: <https://www.510kdatabase.net/k770756/>**SUBMISSION DETAILS**

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|                       |                                    |
|-----------------------|------------------------------------|
| Decision              | Substantially Equivalent (Cleared) |
| Submission type       | Traditional                        |
| Device classification | Echocardiograph (DXK)              |
| Date received         | Apr 25, 1977                       |
| Decision date         | May 9, 1977                        |
| Days to decision      | 14 days                            |
| Third-party review    | No                                 |

**APPLICANT**

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|----------------|---|
| Company        | <b>Honeywell, Inc.</b>                  |
| Location       | Mchenry, IL, US                         |
| 510(k) history | 69 submissions · 69 cleared · 1976-1990 |

Honeywell, Inc. is an American multinational conglomerate headquartered in Charlotte, North Carolina. The company operates across aerospace, building automation, industrial automation, and energy solutions. Honeywell's medical device regulatory history spans from 1976 to 1990. The company received FDA 510(k) clearances from total submissions. Cardiovascular devices represented the dominant focus, accounting for approximately 75% of submissions. This historical record reflects the company's past involvement in patient monitoring systems, defibrillators, and related cardiov...

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Device record: <https://www.510kdatabase.net/k770756/> Data sourced from FDA 510(k) public records ([accessdata.fda.gov](http://accessdata.fda.gov)).

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