

**K760298 SUPPORT, WRIST, FIRM WIRE-FOAM**Sep 8, 1976  
47 days to decisionK760298 · Product code: **ILH** · Physical Medicine  
Source: <https://www.510kdatabase.net/k760298/>**SUBMISSION DETAILS**

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|                       |                                    |
|-----------------------|------------------------------------|
| Decision              | Substantially Equivalent (Cleared) |
| Submission type       | Traditional                        |
| Device classification | Splint, Hand, And Components (ILH) |
| Date received         | Jul 23, 1976                       |
| Decision date         | Sep 8, 1976                        |
| Days to decision      | 47 days                            |
| Third-party review    | No                                 |

**APPLICANT**

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|----------------|---|
| Company        | <b>Fred Sammons, Inc.</b>   |
| Location       | Mchenry, IL, US   |
| Website        | <a href="https://www.sammons-preston.com">https://www.sammons-preston.com</a> |
| 510(k) history | 278 submissions · 278 cleared · 1976-1988                                     |

Fred Sammons, Inc. is a medical device company based in McHenry, US, specializing in rehabilitation products and assistive devices for patient care and therapy. The company has received FDA 510(k) clearances from total submissions, with 88% focused on Physical Medicine devices. FDA 510(k) clearances span from 1976 to 1988, establishing a historical regulatory record in therapeutic aids, orthotic supports, and mobility assistance equipment. Notable cleared devices include orthotic plastics, arm slings, splints, traction exercise equipment, and specialized assessment tools....

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510k Database - [www.510kdatabase.net](http://www.510kdatabase.net)Device record: <https://www.510kdatabase.net/k760298/> Data sourced from FDA 510(k) public records ([accessdata.fda.gov](http://accessdata.fda.gov)).510k Database is an independent platform not affiliated with the U.S. Food and Drug Administration. Always verify at [accessdata.fda.gov](http://accessdata.fda.gov). - Generated May 28, 2026