

K161928 CAPI 3 IMMUNOTYPING, CAPILLARYS 3 TERA, IT/IF CONTROLDec 21, 2016
160 days to decisionK161928 · Product code: **CFF** · Immunology
Source: <https://www.510kdatabase.net/k161928/>**SUBMISSION DETAILS**

| | |
|-----------------------|--|
| Decision | Substantially Equivalent (Cleared) |
| Submission type | Traditional |
| Device classification | Immuno-electrophoretic, Immunoglobulins, (g, A, M) (CFF) |
| Date received | Jul 14, 2016 |
| Decision date | Dec 21, 2016 |
| Days to decision | 160 days |
| Third-party review | No |
| Summary / Statement | Summary |

APPLICANT

| | |
|----------------|---|
| Company | Sebia |
| Location | Chelsea, MI, US |
| Contact | Karen Anderson |
| Website | http://www.sebia.com/ |
| 510(k) history | 32 submissions · 32 cleared · 1995-2024 |

Sebia is a global specialized in vitro diagnostic (IVD) player providing powerful diagnostic tools for chronic and metabolic diseases. The company operates with a manufacturing facility in Chelsea, US, and serves laboratories worldwide with instruments, tests, and software solutions. Sebia has received FDA 510(k) clearances from total submissions since 1995, with no denied submissions on record. The company specializes in immunology devices, including capillary electrophoresis and immunofixation technologies. Latest clearance in 2024 confirms active regulatory engagement....

510k Database - www.510kdatabase.net

Device record: <https://www.510kdatabase.net/k161928/> Data sourced from FDA 510(k) public records (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. - Generated May 14, 2026