

**K110687 RELIEVA STRATUS PRO MICROFLOW SPACER  
(FRONTAL)**Oct 7, 2011  
210 days to decisionK110687 · Product code: **KAM** · General & Plastic Surgery  
Source: <https://www.510kdatabase.net/k110687/>**SUBMISSION DETAILS**

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|                       |                                    |
|-----------------------|------------------------------------|
| Decision              | Substantially Equivalent (Cleared) |
| Submission type       | Special                            |
| Device classification | Cannula, Sinus (KAM)               |
| Date received         | Mar 11, 2011                       |
| Decision date         | Oct 7, 2011                        |
| Days to decision      | 210 days                           |
| Third-party review    | No                                 |
| Summary / Statement   | Summary                            |

**APPLICANT**

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|----------------|---|
| Company        | <b>Acclarent, Inc.</b>  |
| Location       | Irvine, CA, US  |
| Contact        | DEAN KNIGHT   |
| Website        | <a href="https://www.acclarent.com">https://www.acclarent.com</a> |
| 510(k) history | 45 submissions · 44 cleared · 2005-2026                           |

Acclarent, Inc. is a subsidiary of Integra LifeSciences based in Irvine, California. The company develops technology for Ear, Nose, Throat related conditions. Acclarent has received FDA 510(k) clearances from total submissions since its first clearance in 2005. Ear, Nose, Throat devices represent the dominant focus, accounting for 76% of all submissions. The company's latest clearance was in 2026, demonstrating continued regulatory activity. The company specializes in minimally invasive surgical instruments and balloon dilation systems for sinus and Eustachian tube proced...

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510k Database - [www.510kdatabase.net](http://www.510kdatabase.net)

Device record: <https://www.510kdatabase.net/k110687/> Data sourced from FDA 510(k) public records ([accessdata.fda.gov](http://accessdata.fda.gov)).

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