

**K060107 MODIFICATION TO: EXACTECH ZIRAMIC ZIRCONIA  
12/14 FEMORAL HEADS**Feb 6, 2006  
24 days to decisionK060107 · Product code: **LZO** · Orthopedic  
Source: <https://www.510kdatabase.net/k060107/>**SUBMISSION DETAILS**

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|                       |  |
|-----------------------|--|
| Decision              | Substantially Equivalent (Cleared)   |
| Submission type       | Special  |
| Device classification | Prosthesis, Hip, Semi-constrained, Metal/ceramic/polymer, Cemented Or Non-porous, Uncemented (LZO) |
| Date received         | Jan 13, 2006   |
| Decision date         | Feb 6, 2006  |
| Days to decision      | 24 days  |
| Third-party review    | No   |
| Summary / Statement   | Summary  |

**APPLICANT**

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|----------------|---|
| Company        | <b>Exactech, Inc.</b>                                     |
| Location       | Gainesville, FL, US                                       |
| Contact        | MARITZA ELIAS   |
| Website        | <a href="https://www.exac.com/">https://www.exac.com/</a> |
| 510(k) history | 186 submissions · 174 cleared · 1986-2026                 |

Exactech, Inc. operates with a manufacturing facility in Gainesville, US. The company does not offer direct sales or distribution in the United States. Product inquiries and safety concerns are handled through designated company contacts. Exactech has submitted FDA 510(k) applications, resulting in cleared devices. The company's regulatory activity spans from 1986 to 2026, demonstrating sustained engagement with FDA clearance processes. Orthopedic devices represent the dominant focus of the company's portfolio, accounting for approximately 99% of submissions. Recent FDA 5...

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