

**K233630 Ambu® aScope™ 5 Uretero (Standard Deflection)**Jun 24, 2024  
224 days to decisionK233630 · Product code: **FGB** · Gastroenterology & Urology  
Source: <https://www.510kdatabase.net/k233630/>**SUBMISSION DETAILS**

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|                       |   |
|-----------------------|---|
| Decision              | Substantially Equivalent (Cleared)                          |
| Submission type       | Traditional   |
| Device classification | Ureteroscope And Accessories, Flexible/rigid (FGB)          |
| Date received         | Nov 13, 2023  |
| Decision date         | Jun 24, 2024  |
| Days to decision      | 224 days  |
| Third-party review    | No  |
| Combination product   | No  |
| PCCP authorized       | No  |
| Summary / Statement   | Summary   |
| Other names           | Ambu® aScope™ 5 Uretero (Reverse Deflection); Ambu® aBox™ 2 |

**APPLICANT**

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|----------------|---|
| Company        | <b>Ambu A/S</b>   |
| Location       | Glen Burnie, MD, US                                     |
| Contact        | Mingye Zhang  |
| Website        | <a href="https://www.ambu.com">https://www.ambu.com</a> |
| 510(k) history | 38 submissions · 38 cleared · 2005-2026                 |

Ambu A/S is a global medical device company specializing in single-use endoscopy and airway management solutions. The company operates with a manufacturing facility in Glen Burnie, Maryland, and serves hospitals and emergency care settings worldwide. Ambu created the single-use endoscopy market in 2009 and remains the market leader in this category. Ambu has received FDA 510(k) clearances from total submissions, with no denied submissions on record. The company's regulatory activity spans from 2005 to 2026, demonstrating sustained innovation and market presence. Recent cl...

**REGULATORY CONSULTANT**

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|-----------------|-------------------|
| Consulting firm | <b>Ambu, Inc.</b> |
| Contact         | Sanjay Parikh     |

Regulatory consulting firm that managed this 510(k) submission on behalf of the applicant. Source: [FDA accessdata.fda.gov](https://accessdata.fda.gov)

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

Device record: <https://www.510kdatabase.net/k233630/> Data sourced from FDA 510(k) public records ([accessdata.fda.gov](https://accessdata.fda.gov)).

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