

**K972540 S-357 VIDEO IMAGING SYSTEM**Sep 18, 1997  
73 days to decisionK972540 · Product code: **FET** · Gastroenterology & Urology  
Source: <https://www.510kdatabase.net/k972540/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Endoscopic Video Imaging System/component, Gastroenterology-urology (FET)
Date received	Jul 7, 1997
Decision date	Sep 18, 1997
Days to decision	73 days
Third-party review	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Sopro</b>
Location	Marseille, FR
Contact	PIERRE MONTILLOT
Website	<a href="http://www.soprole.cl/">http://www.soprole.cl/</a>
510(k) history	25 submissions · 25 cleared · 1997-2023

Sopro specialized in surgical visualization and imaging devices for general and plastic surgery applications. The company operated a manufacturing facility in Marseille, France. Sopro received FDA 510(k) clearances from total submissions between 1997 and 2023. The company focused exclusively on General & Plastic Surgery devices, particularly endoscopy cameras, digital operating room cameras, and light source systems. All submissions resulted in clearance with no denials on record. The company's product portfolio included digital endoscopy cameras, laparoscopes, operating ...

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