

**K093622 AEM MONITORING SYSTEM, MODELS EM2, EMR, ES4007, ES4107, ES9005 AND ES9015**Aug 20, 2010  
270 days to decisionK093622 · Product code: **GEI** · General & Plastic Surgery  
Source: <https://www.510kdatabase.net/k093622/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Electrosurgical, Cutting & Coagulation & Accessories (GEI)
Date received	Nov 23, 2009
Decision date	Aug 20, 2010
Days to decision	270 days
Third-party review	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Encision, Inc.</b>
Location	Boulder, CO, US
Contact	JUDITH V KING
Website	<a href="http://encision.com/">http://encision.com/</a>
510(k) history	10 submissions · 10 cleared · 2007-2020

Encision, Inc. designs and manufactures laparoscopic surgical instruments featuring Active Electrode Monitoring (AEM®) technology. The company, with a manufacturing facility in Boulder, US, specializes in burn protection systems and shielded surgical instruments for minimally invasive procedures. Encision has received FDA 510(k) clearances from total submissions since 2007. All cleared devices fall within the General & Plastic Surgery category. The company's last FDA 510(k) clearance was in 2020, and the company is currently inactive with no recent submissions. The compan...

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

Device record: <https://www.510kdatabase.net/k093622/> Data sourced from FDA 510(k) public records ([accessdata.fda.gov](http://accessdata.fda.gov)).

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