

**K915058 HEMAGEN ENA SCREENING KIT (EIA METHOD)**Dec 20, 1991  
42 days to decisionK915058 · Product code: LLL · Immunology  
Source: <https://www.510kdatabase.net/k915058/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Extractable Antinuclear Antibody, Antigen And Control (LLL)
Date received	Nov 8, 1991
Decision date	Dec 20, 1991
Days to decision	42 days
Third-party review	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Hemagen Diagnostics, Inc.</b>
Location	Waltham, MA, US
Contact	A WILLAND
Website	<a href="http://www.hemagen.com/">http://www.hemagen.com/</a>
510(k) history	52 submissions · 52 cleared · 1986-2004

Hemagen Diagnostics, Inc. was founded in 1985 by scientists from Boston University School of Medicine. The company provides clinical diagnostic solutions specializing in immunology devices for autoimmune and infectious disease testing. Hemagen offers gold standard IFA products, ELISA, HA, and point-of-care testing formats for human and veterinary diagnostics. Hemagen has received FDA 510(k) clearances from total submissions since its first clearance in 1986. The company's regulatory portfolio focuses on immunology devices, including antibody detection kits, autoimmune scr...

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