

K925425 SPECIFIC IGE EIA-TURBO ASSAYJan 14, 1993
79 days to decisionK925425 · Product code: **DHB** · Immunology
Source: <https://www.510kdatabase.net/k925425/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	System, Test, Radioallergosorbent (rast) Immunological (DHB)
Date received	Oct 27, 1992
Decision date	Jan 14, 1993
Days to decision	79 days
Third-party review	No
Summary / Statement	Statement

APPLICANT

Company	Hycor Biomedical, Inc.
Location	Garden Grove, CA, US
Contact	Anne Jepson
510(k) history	51 submissions · 51 cleared · 1989-2008

Hycor Biomedical, Inc. is an American manufacturer of in vitro diagnostic products for blood testing. The company is based in Garden Grove, California and specializes in allergy and autoimmune testing solutions. Hycor Biomedical has received FDA 510(k) clearances from total submissions since its first clearance in 1989. The company's regulatory portfolio is dominated by Immunology devices, which represent the majority of its cleared submissions. The latest FDA 510(k) clearance on record dates to 2008, reflecting the company's historical significance in the diagnostic devi...

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