

K931620 OPTI-FREE CONTACT LENS CASEJun 28, 1993
87 days to decisionK931620 · Product code: **LRX** · Ophthalmic
Source: <https://www.510kdatabase.net/k931620/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Case, Contact Lens (LRX)
Date received	Apr 2, 1993
Decision date	Jun 28, 1993
Days to decision	87 days
Third-party review	No
Summary / Statement	Statement

APPLICANT

Company	Alcon Laboratories
Location	Mchenry, IL, US
Contact	DEAN A KNIGHT
Website	https://www.alcon.com
510(k) history	47 submissions · 47 cleared · 1976-2007

Alcon Laboratories is a Swiss-American pharmaceutical and medical device company specializing in eye care products. The company maintains operational headquarters in Fort Worth, Texas, with a significant global presence in eye care innovation. Alcon has received FDA 510(k) clearances from total submissions since its first clearance in 1976. The company's regulatory portfolio is dominated by Ophthalmic devices, which account for 74% of all submissions. The latest clearance on record dates to 2007, reflecting the company's historical significance in the medical device regul...
