

K013232 CYBERCASES BY BAUSCH & LOMBNov 20, 2001
54 days to decisionK013232 · Product code: **LRX** · Ophthalmic
Source: <https://www.510kdatabase.net/k013232/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Case, Contact Lens (LRX)
Date received	Sep 27, 2001
Decision date	Nov 20, 2001
Days to decision	54 days
Third-party review	No
Summary / Statement	Summary

APPLICANT

Company	Bausch & Lomb, Inc.
Location	Mchenry, IL, US
Contact	KIM S DEVITTO
Website	http://www.bausch.com
510(k) history	92 submissions · 92 cleared · 1977-2019

Bausch & Lomb, Inc. is a Canadian eye health company founded in 1853. The company is now part of Valeant Pharmaceuticals following a 2013 acquisition. Bausch & Lomb has received FDA 510(k) clearances from total submissions since 1977. The company specializes in Ophthalmic devices, which represent 83% of its regulatory submissions. Recent cleared devices include contact lenses, intraocular lens injectors, lens delivery systems, and care solutions. The company's last FDA 510(k) clearance was in 2019, and this profile reflects its historical regulatory record. Bausch & Lomb ...
