

**K983150 COMPLETE BRAND LUBRICATING AND REWETTING DROPS**Nov 25, 1998  
77 days to decisionK983150 · Product code: **LPN** · Ophthalmic  
Source: <https://www.510kdatabase.net/k983150/>**SUBMISSION DETAILS**

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Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Accessories, Soft Lens Products (LPN)
Date received	Sep 9, 1998
Decision date	Nov 25, 1998
Days to decision	77 days
Third-party review	No
Summary / Statement	Summary

**APPLICANT**

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Company	<b>Allergan, Inc.</b>
Location	Walker, MI, US
Contact	PAUL J NOWACKI
Website	<a href="http://www.allergan.com">http://www.allergan.com</a>
510(k) history	33 submissions · 33 cleared · 1982-2019

Allergan, Inc. was an American global pharmaceutical company headquartered in Walker, US. The company focused on eye care, neurosciences, medical dermatology, and medical aesthetics before ceasing independent operations in 2015. Allergan received FDA 510(k) clearances from total submissions between 1982 and 2019. The company's regulatory portfolio was dominated by Ophthalmic devices, which accounted for 88% of all submissions. Notable cleared products include the XEN Glaucoma Treatment System, REFRESH rewetting drops, and the Natrelle tissue expander for surgical applicat...

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Device record: <https://www.510kdatabase.net/k983150/> Data sourced from FDA 510(k) public records ([accessdata.fda.gov](http://accessdata.fda.gov)).

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