

## References for CEQUA® (cyclosporine ophthalmic solution) 0.09%

**Phase 4 Study Design:** Single arm, Phase 4, 12-week, multicenter study of 124 adults with DED inadequately controlled (ie, still symptomatic and/or exhibiting disease signs) on current Restasis® therapy. The co-primary endpoints were corneal fluorescein staining (CFS) and modified Symptom Assessment in Dry Eye (mSANDE) at Week 12. Patients received 1 drop, 2X daily of CEQUA in each eye.<sup>1,2,5</sup>

Enrolled patients were selected by their doctors based on: Clinical diagnosis of DED and treatment on Restasis for  $\geq 3$  months; BCVA of  $\geq 20/200$ ; mSANDE score of  $\geq 40$ ; total CFS  $\geq 6$  or CFS in an individual zone  $\geq 2$  at baseline.<sup>5</sup>

Exclusions: Previous history of failure on Restasis; discontinued/switched to a different immunomodulatory; allergic conjunctivitis; stable dose for  $\geq 3$  months of immunomodulators, antihistamines, cholinergics, antimuscarinics, phenothiazines, retinoids, or any systemic or topical corticosteroids.<sup>1,5</sup>

### References:

1. Data on file. Cranbury, NJ: Sun Pharmaceutical Industries, Inc.
2. Johnston, J. Effect of OTX-101 0.09% on corneal staining and SANDE scores in patients with dry eye disease uncontrolled on cyclosporine ophthalmic emulsion 0.05%. Abstract presented at American Academy of Optometry 2023; October 12, 2023; New Orleans, LA.
3. CEQUA [package insert]. Cranbury, NJ: Sun Pharmaceutical Industries, Inc.; 2022.
4. Goldberg DF, Malhotra RP, Schechter BA, Justice A, Weiss SL, Sheppard JD. A phase 3, randomized, double-masked study of OTX-101 ophthalmic solution 0.09% in the treatment of dry eye disease. *Ophthalmology*. 2019;126(9):1230-1237.
5. Effect of CEQUA in Subjects with Dry Eye Disease, ClinicalTrials.gov identifier NCT04357795. Updated Sept 09, 2022. Accessed August 29, 2023. <https://www.clinicaltrials.gov/study/NCT04357795>

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