

### UnitedHealthcare Pharmacy Clinical Pharmacy Programs

| Program Number    | 2023 1088-14   |
|-------------------|--|
| Program           | Prior Authorization-Notification   |
| Medication        | Cequa <sup>™</sup> (cyclosporine 0.09% ophthalmic solution)*, Miebo <sup>™</sup> |
|                   | (perfluorohexyloctane)*, Restasis <sup>®</sup> (cyclosporine 0.05% ophthalmic    |
|                   | emulsion), Restasis MultiDose <sup>™</sup> (cyclosporine 0.05% ophthalmic        |
|                   | emulsion)*, Tyrvaya <sup>™</sup> (varenicline nasal spray), Vevye <sup>™</sup>   |
|                   | (cyclosporine 0.1%)*, Xiidra® (lifitegrast 5% ophthalmic solution)               |
| P&T Approval Date | 3/2006, 3/2007, 8/2008, 8/2009, 9/2010, 3/2011, 2/2012, 2/2013,                  |
|                   | 4/2014, 4/2015, 3/2016, 12/2016, 9/2017, 9/2018, 3/2019, 4/2020,                 |
|                   | 4/2021, 12/2021, 12/2022, 9/2023   |
| Effective Date    | 12/1/2023  |

## 1. Background:

Cequa (cyclosporine 0.09% ophthalmic solution)\*, Restasis (cyclosporine 0.05% ophthalmic emulsion) and Restasis MultiDose (cyclosporine 0.05% ophthalmic emulsion)\*, are indicated to increase tear production in patients whose tear production is presumed to be suppressed due to ocular inflammation associated with keratoconjunctivitis sicca.

Miebo (perfluorohexyloctane)\*, Tyrvaya (varenicline nasal spray), Vevye (cyclosporine 0.1%)\* and Xiidra (lifitegrast 5% ophthalmic solution) are indicated for the treatment of the signs and symptoms of dry eye disease.

#### 2. Coverage Criteria<sup>a</sup>:

# A. Initial Authorization

- 1. Cequa\*, Miebo\*, Restasis, Restasis MultiDose\*, Tyrvaya, Vevye\* or Xiidra will be approved based on the following criterion:
  - a. Diagnosis of **<u>one</u>** of the following:
    - 1) Moderate to severe keratoconjuctivitis sicca
    - 2) Dry Eye Disease

#### Authorization will be issued for 6 months.

#### B. <u>Reauthorization</u>

- 1. Cequa\*, Miebo\*, Restasis, Restasis MultiDose\*, Tyrvaya, Vevye\* or Xiidra will be approved based on the following criterion:
  - a. Patient has demonstrated clinically significant improvement with therapy

# Authorization will be issued for 12 months.

<sup>a</sup> State mandates may apply. Any federal regulatory requirements and the member specific benefit plan coverage may also impact coverage criteria. Other policies and utilization management programs may apply.

\*Cequa, Miebo, Restasis MultiDose and Vevyeare typically excluded from coverage.



# 3. Additional Clinical Rules:

- Notwithstanding Coverage Criteria, UnitedHealthcare may approve initial and re-authorization based solely on previous claim/medication history, diagnosis codes (ICD-10) and/or claim logic. Use of automated approval and re-approval processes varies by program and/or therapeutic class.
- Supply limits may be in place.
- Prior Authorization Medical Necessity may be in place.

### 4. References:

- 1. Cequa [package insert]. Cranbury, NJ: Sun Pharmaceutical Industries, Inc; December 2022
- 2. Miebo [package insert]. Bridgewater, NJ: Bausch & Lomb Americas Inc; May 2023.
- 3. Restasis [package insert]. Irvine CA: Allergan, Inc.: July 2017.
- 4. Restasis MultiDose [package insert]. Irvine, CA: Allergan, Inc.; October 2016.
- 5. Tyrvaya [package insert]. Princeton NJ: Oyster Point Pharma, Inc; October 2021.
- 6. Vevye [package insert]. Irvine CA: Alliance Medical Products, Inc; June 2023.
- 7. Xiidra [package insert]. East Hanover, NJ: Novartis Pharmaceuticals Corporation; June 2020.
- 6. American Academy of Ophthalmology. Dry Eye Syndrome Preferred Practice Pattern 2018. Accessed October 2022

| Program        | Notification – Dry Eye Disease  |
|----------------|---|
| Change Control |   |
| 4/2014         | No changes to criteria. Updated references.                               |
| 4/2015         | Updating authorization criteria to 6 months and reauthorization criteria  |
|                | to 12 months to align with prior authorization-medical necessity criteria |
| 3/2016         | Removed the Prior Authorization-Medical Necessity program language.       |
| 12/2016        | Updated criteria to allow for Dry Eye Disease. Added Xiidra to criteria.  |
|                | Changed name of criteria to Dry Eye Disease.                              |
| 9/2017         | Annual review. Administrative updates. Added Restasis MultiDose.          |
|                | Updated references.   |
| 9/2018         | Annual review. Administrative updates and updated references.             |
| 12/2018        | Administrative change to add statement regarding use of automated         |
|                | processes.  |
| 3/2019         | Added Cequa and updated references.                                       |
| 4/2020         | Annual review. Removed extemporaneously compounded cyclosporine           |
|                | criteria. Updated references.   |
| 4/2021         | Annual review. Updated references.  |
| 12/2021        | Added Tyrvaya.  |
| 12/2022        | Annual review. Added state mandate language. Updated references.          |
| 9/2023         | Added Miebo and Vevye. Updated references.                                |