# **CLINICAL POLICY**Quantity Limit Override



# Clinical Policy: Quantity Limit Override and Dose Optimization

Reference Number: PA.CP.PMN.59

Effective Date: 01/2018 Last Review Date: 07/2023

**Revision Log** 

# **Description**

This policy establishes the criteria for overriding set quantity limits (QL) and dose optimization.

# **FDA Approved Indication(s)**

Varies by drug product.

#### Policy/Criteria

\* Provider <u>must</u> submit documentation (including office chart notes and lab results) supporting that member has met all approval criteria \*

It is the policy of PA Health & Wellness <sup>®</sup> that dose optimization is implemented when clinically appropriate. Prescribers are required to consolidate multiple units of lower strength to the fewest units required to achieve the desired daily dose/regimen based on commercially available dosage strengths (see *Appendix D* for examples). Requests for multiple units of a lower strength will be denied when the plan-approved QL for such medication is exceeded and higher strength units are commercially available to achieve the desired daily dose/regimen.

It is the policy of PA Health & Wellness® that quantity limit (QL) and dose optimization exceptions are **medically necessary** when the following criteria are met:

# I. Initial Approval Criteria

#### A. Quantity Limit Exceptions (must meet all):

Refer to Section IB for conditions eligible for continuity of care and Section IC for pain management

- 1. One of the following (a or b):
  - a. Requested dose is supported by practice guidelines or peer-reviewed literature (e.g., phase 3 study or equivalent published in a reputable peer reviewed medical journal or text) for the relevant off-label use and/or regimen (*prescriber must submit supporting evidence*);
  - b. Diagnosis of a condition/disease for which FDA dosing guidelines indicate a higher quantity (dose or frequency) than the currently set QL; Example: Proton pump inhibitors, which are commonly used for gastroesophageal reflux disease, have a QL of one dose per day; however, when there is a rare diagnosis such as Zollinger-Ellison syndrome, an override for two doses per day is allowed
- 2. One of the following (a or b):
  - a. Member has been titrated up from the lower dose with partial improvement without adverse reactions;
  - b. For retail fill quantity limit, quantity limit requires multiple trips to the pharmacy in one month to complete therapy;
- 3. Dose optimization is required, unless one of the following applies (a or b):

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- a. Dose titration: Multiple lower strength units are requested for the purpose of dose titration:
- b. Other clinical reasons: Medical justification supports inability to use the higher strength units to achieve the desired dose/regimen.

Approval duration: 12 months (60 days if dose optimization exception is requested due to dose titration)

# **B.** Continuity of care:

1. Currently receiving medication via PA Health & Wellness benefit and documentation supports positive response to therapy or the Continuity of Care policy. Refer to the Continuity of Care Policy (PA.LTSS.PHARM.01).

# C. Narcotic Analgesics QL Exceptions

- 1. Refer to Long-Acting Opioid Analgesics policy, PHW.PDL.110
- 2. Refer to Short-Acting Opioid Analgesics policy, PHW.PDL.109

# **D. Dose Optimization Exceptions** (must meet all):

- 1. One of the following (a or b):
  - a. Dose titration: Multiple lower strength units are requested for the purpose of dose titration;
  - b. Other clinical reasons: Medical justification supports inability to use the higher strength units to achieve the desired dose/regimen;
- 2. Request meets one of the following (a or b):
  - a. Dose does not exceed the FDA-recommended regimen and maximum daily dose;
  - b. Requested dose is supported by practice guidelines or peer-reviewed literature for the relevant off-label use (*prescriber must submit supporting evidence*).

#### **Approval duration:**

**Dose titration** – Duration of request or 60 days, whichever is less **Other clinical reasons** – Duration of request or 12 months, whichever is less

#### **II. Continued Therapy:**

## **A. All Requests in Section I** (must meet all):

- 1. Currently receiving medication via PA Health & Wellness benefit and documentation supports positive response to therapy;
- 2. Dose optimization is required, unless one of the following applies (a or b):
  - a. Documentation supports the continued need for dose titration or tapering;
  - b. Medical justification supports inability to use the higher strength units to achieve the desired dose/regimen;
- 3. Request meets one of the following (a or b):
  - a. Dose does not exceed the FDA-recommended regimen and maximum daily dose;
  - b. Requested dose is supported by practice guidelines or peer-reviewed literature for the relevant off-label use (*prescriber must submit supporting evidence*).

Approval duration: 12 months (60 days if dose optimization exception is requested due to dose titration)



# III. Diagnoses/Indications for which coverage is NOT authorized: Not Applicable

# IV. Appendices/General Information

Appendix A: Abbreviation Key

FDA: Food and Drug Administration

QL: quantity limit

Appendix B: Therapeutic Alternatives

Not applicable

Appendix C: Contraindications/Boxed Warnings

Varies by drug product

# Appendix D: General Information

• Dose optimization is the consolidation of multiple units of lower strength to the fewest units required to achieve the desired daily dose/regimen based on commercially available dosage strengths. This can reduce pill burden, simplify therapeutic regimens, improve treatment compliance, and reduce pharmacy spend. Requests for multiple units of a lower strength will be denied when the plan-approved QL for such medication is exceeded and higher strength units are commercially available to achieve the desired daily dose/regimen.

Request ExamplePrescribed RegimenApprovable RegimenRequest for Seroquel XRSeroquel XR 200 mgSeroquel XR 400 mg800 mg/daytablets, 4 tablets/daytablets, 2 tablets/dayRequest for aripiprazole 30Aripiprazole 15 mg tablets, 2 tablets/dayAripiprazole 30 mgmg/day2 tablets/daytablet, 1 tablet/day

## V. Dosage and Administration

Not applicable

#### VI. Product Availability

Not applicable

#### VII. References

Not applicable

Reviews, Revisions, and Approvals	Date	P&T Approval Date
4Q 2018 annual review: converted to new template; combined criteria sets for rare conditions and off-label use to apply more broadly; added oncology to list of possible continuation of care eligible conditions; referred off-label dosing to the off-label use policy; references reviewed and updated.	08/2018	

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Reviews, Revisions, and Approvals	Date	P&T Approval Date
09/01/2019 statewide PDL submission: Revised references to	09/2019	
Long- and short-acting opioid policies to account for name change		
occurring with statewide PDL implementation 01/01/2020;		
removed reference to PA.CP.PMN.53		
4Q 2020 annual review: References reviewed and updated.	07/2020	
4Q 2021 annual review: no significant changes	10/2021	
Added dose optimization criteria (PA.CP.PMN.13 retired).	07/2022	
4Q 2022 annual review: no significant changes.	10/2022	
Added retail fill limit override path	07/2023	