



SmartPA Criteria Proposal

Drug/Drug Class:	Sympatholytics PDL Edit		
First Implementation Date:	January 5, 2017		
Proposed Date:	September 15, 2022		
Prepared For:	MO HealthNet		
Prepared By:	MO HealthNet/Conduent		
Criteria Status:	□ Existing Criteria⋈ Revision of Existing Criteria□ New Criteria		

Executive Summary

Purpose: The MO HealthNet Pharmacy Program will implement a state-specific preferred drug list.

Why Issue Selected:

The sympatholytic agents are indicated for various functions for example, treatment of hypertension, anxiety, panic disorder, post-traumatic stress disorder and withdrawal symptoms from opioids. The oral, centrally-acting, alpha-2 adrenergic receptor agonists include clonidine and guanfacine. Methyldopa continues to be used to treat hypertension in pregnant women. A diuretic, usually hydrochlorothiazide or chlorthalidone may be combined with clonidine or methyldopa. The sympatholytics are typically part of a multiple antihypertensive drug regimen for participants who have not reached their target blood pressure. Clonidine is also available in a transdermal formulation for the treatment of hypertension.

Total program savings for the PDL classes will be regularly reviewed.

Program-	Preferred Agents	Non-Preferred Agents		
Specific	Clonidine Patch	Catapres® Tabs		
Information:	Clonidine Tabs	Catapres-TTS® Patch		
	Guanfacine	Lucemyra®		
	 Methyldopa 	Methyldopa/HCTZ		
Type of Criteria:	☐ Increased risk of ADE	☑ Preferred Drug List		
		☐ Clinical Edit		
Data Sources:	☐ Only Administrative Databases	☑ Databases + Prescriber-Supplied		

Setting & Population

- Drug class for review: Sympatholytics
- Age range: All appropriate MO HealthNet participants

Approval Criteria

- Failure to achieve desired therapeutic outcomes with trial on 3 or more preferred agents:
 - o Documented trial period for preferred agents **OR**
 - Documented ADE/ADR to preferred agents
- For Lucemyra:
 - o Documented diagnosis of opioid withdrawal AND
 - o Documented trial period of clonidine oral OR patch

Е		Crite	

 Lack of adequate trial on required preferred agents Therapy will be denied if all approval criteria are not met
Required Documentation
Laboratory Results: Progress Notes: Other:
Disposition of Edit
Denial: Exception Code "0160" (Preferred Drug List) Rule Type: PDL
Default Approval Period
1 year

References

- Evidence-Based Medicine Analysis: "Sympatholytic Antihypertensives", UMKC-DIC; July 2022.
- Evidence-Based Medicine and Fiscal Analysis: "Sympatholytic Antihypertensive Agents Therapeutic Class Review", Conduent Business Services, L.L.C., Richmond, VA; July 2021.
- USPDI, Micromedex; 2022.
- Facts and Comparisons eAnswers (online); 2022 Clinical Drug Information, LLC.