



# SmartPA Criteria Proposal

Drug/Drug Class:	Glucagon-Like Peptide -1 (GLP-1) Receptor Agonists & Combination Agents PDL Edit	
First Implementation Date:	October 7, 2010	
Revised Date:	October 1, 2020	
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:

Type 2 diabetes mellitus is a significant health problem associated with excessive morbidity and mortality. As the prevalence of this metabolic disorder is rapidly increasing and as older treatments fail to stabilize the disease in many participants, prevention and control are considered key objectives. Metformin is still the cornerstone of type 2 diabetes mellitus treatment however many patients will require an additional agent(s). According to the ADA, several classes can be considered as add-on therapy, including the glucagon-like peptide-1 (GLP-1) receptor agonists. Selection of a specific agent should be based on drug-specific characteristics (e.g., adverse events, weight gain, hypoglycemia risk, cost) and patient preferences. Based on differences in cardiovascular risk/benefit and weight gain among the GLP-1 receptor agonists, patients with certain compelling indications might benefit from a specific agent in the class. For patients with established atherosclerotic cardiovascular disease, Victoza® (liraglutide), Trulicity® (dulaglutide) and injectable Ozempic® (semaglutide) have all demonstrated cardiovascular benefit are preferred, as they are FDA-approved for cardiovascular disease reduction. For patients with a compelling need for weight loss, semaglutide is associated with the largest weight reduction. GLP-1 receptor agonists have a similar safety profile with gastrointestinal disorders being the most common adverse effect. All GLP-1 receptor agonists, except Byetta® (exenatide) and Adlyxin™ (lixisenatide), have a boxed warning regarding the risk of thyroid tumors. Dual therapy with insulin and a GLP-1 receptor agonist can be considered if patients cannot meet their HbA1c goals with basal insulin or a GLP-1 receptor agonist alone. No significant efficacy or safety differences have been noted between Xultophy® (insulin degludec/liraglutide) and Soliqua® (insulin glargine/lixisenatide).

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

Program-Specific	Preferred Agents	Non-Preferred Agents	
Information:	Bydureon®     Ryotto®	Adlyxin™     Rydureon® Reise™ Auto Injector	
	Byetta®     Victoza®	<ul> <li>Bydureon<sup>®</sup> Bcise<sup>™</sup> Auto Injector</li> <li>Ozempic<sup>®</sup></li> </ul>	
	V101024	Rybelsus®	
		• Soliqua®	
		Trulicity®	
		Xultophy®	
Type of Criteria:	☐ Increased risk of ADE	☑ Preferred Drug List	
••	☐ Appropriate Indications	☐ Clinical Edit	
Data Sources:	□ Only Administrative Databases	M Databases + Proscriber Supplied	
Data Sources.	☐ Only Administrative Databases	□ Databases + Prescriber-Supplied	
Setting & Popula	ation		
<ul> <li>Drug class for review: Glucagon-Like Peptide -1 (GLP-1) Receptor Agonists &amp; Combination Agents</li> <li>Age range: All appropriate MO HealthNet participants aged 10 years and older</li> </ul>			
Age range. An	appropriate ino riealtinet participants age	a To years and older	
Approval Criteria			
Participants aged 18 years or older AND			
<ul> <li>Adequate therapeutic trial of metformin in the past year AND</li> <li>Failure to achieve desired therapeutic outcomes with trial on 2 or more preferred agents</li> </ul>			
Documented trial period of preferred agents			
<ul> <li>Documented ADE/ADR to preferred agents AND</li> </ul>			
<ul> <li>For Rybelsus: documented therapeutic trial of Ozempic in the past year OR</li> <li>For Victoza: participants aged 10 years or older OR</li> </ul>			
-		trial on 2 or more preferred long acting	
insulins		and on a or more presented teng dealing	
Denial Criteria			
• Lack of adogu	ata trial an required professed agents		
<ul> <li>Lack of adequate trial on required preferred agents</li> <li>Therapy will be denied if no approval criteria are met</li> </ul>			
For exenatide: documented diagnosis of End Stage Renal Disease (ESRD) or severe renal			
	reatinine clearance <30 ml/min)		
Claim exceeds	s maximum dosing limitation for the following	g:	
VICTOZA 18	MG/3 ML PEN LIRAGLUTIDE 0	.3mL per day	
Required Docum	nentation		
Laboratory Resul	ts: Progress Notes:		

## Disposition of Edit

MedWatch Form:

Denial: Exception Code "0160" (Preferred Drug List)

Other:

SmartPA PDL Proposal Form

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Rule Type: PDL

### **Default Approval Period**

1 year

#### References

- 1. Drug Effectiveness Review Project Drug Class Review Newer Diabetes Medications and Combinations. Center for Evidence-Based Policy, Oregon Health & Science University; February 2011; updated July 2016.
- 2. Evidence-Based Medicine and Fiscal Analysis: "GLP-1 Receptor Agonists and Combinations Therapeutic Class Review", Conduent Business Services, L.L.C., Richmond, VA; April 2020.
- 3. Evidence-Based Medicine Analysis: "Antidiabetic Mimetics (GLP-1 Receptor Agonist)", UMKC-DIC; March 2020.
- 4. Evidence-Based Medicine Analysis: "Antidiabetic Combination Agents Oral and Injectable", UMKC-DIC; March 2020.
- 5. American Diabetes Association (ADA). Standards of Medical Care in Diabetes-2020. *Diabetes Care*. 2020;43(suppl 1): S1-S212.
- 6. Lippincott, Williams, Wilkins. PDR Electronic Library, Montvale NJ; 2020.
- 7. USPDI, Micromedex; 2020.
- 8. Facts and Comparisons eAnswers (online); 2020 Clinical Drug Information, LLC.