



# SmartPA Criteria Proposal

Drug/Drug Class:	Isturisa Clinical Edit
First Implementation Date:	TBD
Proposed Date:	December 17, 2020
Prepared for:	MO HealthNet
Prepared by:	MO HealthNet/Conduent
Criteria Status:	□Existing Criteria □Revision of Existing Criteria ⊠New Criteria

## **Executive Summary**

Purpose: Ensure appropriate utilization and control of Isturisa® (osilodrostat)

Why Issue Selected:

Isturisa® (osilodrostat) was FDA approved in March 2020, for the treatment of adult patients with Cushing's disease for whom pituitary surgery is not an option or has not been curative. Cushing's disease occurs in the presence of pituitary adenomas that produce high levels of adrenocorticotropin hormone (ACTH); this oversecretion of ACTH leads to an overproduction of cortisol by the adrenal glands. Cushing's disease can be life threatening if not treated and may cause significant health issues such as obesity, type 2 diabetes, high blood pressure, blood clots, bone loss, immunosuppression, and neuropsychiatric symptoms. Isturisa is a twice-daily oral cortisol synthesis inhibitor that inhibits 11-beta-hydroxylase (CYP11B1) which is responsible for the final step of cortisol biosynthesis in the adrenal gland. Isturisa represents the first agent with this mechanism of action to be approved by the FDA for the treatment of Cushing's disease. Due to the high cost and specific approved indications, MO HealthNet will impose clinical criteria to ensure appropriate utilization of Isturisa.

## Program-Specific Information:

Date Range FFS 10-01-2019 to 9-30-2020				
Drug	Claims	Cost per tablet	Cost per month (MAC)	
ISTURISA 1 MG TAB	0	\$110.00 MAC	Initial dose (2 mg BID): \$13,200	
ISTURISA 5 MG TAB	0	\$400.00 MAC	Avg maint dose (7 mg BID): \$37,200	
ISTURISA 10 MG TAB	0	\$475.00 MAC	Max dose (30 mg BID): \$85,500	

Type of Criteria: ☐ Increased risk of ADE ☐ Preferred Drug List

Data Sources: ☐ Only Administrative Databases ☐ Databases + Prescriber-Supplied

#### **Setting & Population**

- Drug class for review: Isturisa® (osilodrostat)
- Age range: All appropriate MO HealthNet participants aged 18 years or older

#### SmartPA Clinical Proposal Form

## **Approval Criteria**

#### **Initial Therapy:**

- Participant aged 18 years or older AND
- Prescribed by or in consultation with an endocrinologist or other appropriate specialist for the treated disease state AND
- Documented diagnosis of Cushing's disease AND
- Documentation of failed pituitary surgery or contraindication to pituitary surgery AND
- Documentation of baseline electrocardiogram AND
- Documentation of baseline potassium, magnesium, and cortisol levels

#### Continuation of Therapy:

- Initial approval is for 3 months, renewal of prior authorization may be given for up to 12 months
  following documentation of the following along with an expectation of monitoring of potassium and
  magnesium levels:
  - Recent (within the past 3 months) cortisol levels demonstrating mean urine free cortisol (mUFC) ≤ ULN AND
  - Recent (within the past 3 months) electrocardiogram

Denial Criteria		
Therapy will be den	ied if all approval criteria are not met	
<b>Required Documents</b>	ation	
Laboratory Results:	Progress Notes:	

#### Disposition of Edit

Denial: Exception code "0682" (Clinical Edit)

Rule Type: CE

## **Default Approval Period**

3 months

#### References

- Isturisa (osilodrostat) [package insert]. Lebanon, NJ: Recordati Rare Disease Inc; March 2020.
- IPD Analytics. New Drug Review: Isturisa (osilodrostat). March 2020.
- Nieman L., Biller B., Findling J., et al. Treatment of Cushing's Syndrome: An Endocrine Society Clinical Practice Guideline. The Journal of Clinical Endocrinology & Metabolism; 2015;100(8):2807-283.