

PHARMACY POLICY STATEMENT Ohio Medicaid	
DRUG NAME	Saizen (somatropin)
BILLING CODE	Must use valid NDC code
BENEFIT TYPE	Pharmacy
SITE OF SERVICE ALLOWED	Home
COVERAGE REQUIREMENTS	Prior Authorization Required (Non-Preferred Product)
	QUANTITY LIMIT— per diagnosis, see <b>Dosage allowed</b>
LIST OF DIAGNOSES CONSIDERED NOT	Click Here
MEDICALLY NECESSARY	

Saizen (somatropin) is a **non-preferred** product and will only be considered for coverage under the **pharmacy** benefit when the following criteria are met:

Members must be clinically diagnosed with one of the following disease states and meet their individual criteria as stated.

## Adult GROWTH HORMONE DEFICIENCY (GHD) - Adult or Childhood Onset

For *initial* authorization:

- 1. Member must have a documented 90-day trial and failure of Omnitrope 5.8 mg vial; AND
- 2. Member is 18 years of age or older; AND
- 3. Medication must be prescribed by an endocrinologist; AND
- 4. Member must have a diagnosis of GHD confirmed by **one** of the following:
  - a) Chart notes documentation of acquired structural abnormality (*see Appendix*) of the hypothalamus or pituitary and ≥ 3 documented pituitary hormone deficiencies (*see Appendix*) with included lab results and reference ranges;
  - b) Documented childhood-onset of GHD with a documented congenital abnormality (*see Appendix*) of the hypothalamus or pituitary;
  - c) Two pre-treatment peak serum growth hormone (GH) concentration < 5 ng/mL by stimulation testing with included lab results and reference ranges, unless Macrilen (prior authorization required) was used, in which case a GH level must be < 2.8 ng/ml.
- Dosage allowed: Weight based dosing: 0.005 mg/kg/day initially; can be increased as tolerated to not more than 0.01 mg/kg/day after 4 weeks. Non-weight based dosing: starting dose 0.2 mg/day (0.15-0.30 mg/day) and increased every 1-2 months in increments of 0.1-0.2 mg/day, doses vary considerably.

### *If member meets all the requirements listed above, the medication will be approved for 12 months.* For <u>reauthorization</u>:

- 1. Member must be in compliance with all of the initial criteria; AND
- 2. Member's current IGF-1 level not elevated for age/gender (does not apply to members w/ structural abnormality of hypothalamus/pituitary and at least pituitary hormone deficiencies or childhood onset GHD and congenital abnormality of hypothalamus/pituitary).

*If member meets all the reauthorization requirements above, the medication will be approved for an additional 12 months.* 



## Pediatric GROWTH HORMONE DEFICIENCY (GHD)

For *initial* authorization:

- 1. Member must have a documented 90-day trial and failure of Omnitrope 5.8 mg vial; AND
- 2. Member is 17 years old or younger; AND
- 3. Medication must be prescribed by an endocrinologist; AND
- 4. Member must have a diagnosis of GHD confirmed by **one** of the following:
  - a) Neonate or diagnosed with GHD as neonate indicated by ALL of the following:
    - i) Chart notes, labs, and documentation must be included to support the diagnosis (e.g, hypoglycemia with random GH level ≤ 5 ng/mL, evidence of multiple pituitary hormone deficiency (see Appendix), MRI results);
    - ii) Pituitary abnormality (secondary to congenital anomaly (*see Appendix*), pituitary tumor, or irradiation);
    - iii) A known deficiency of at least one other pituitary hormone (see Appendix);
  - b) Two pre-treatment peak serum growth hormone concentration < 10 ng/mL by stimulation testing (*must include lab results with reference ranges*);
  - c) A documented pituitary or CNS disorder and a pre-treatment IGF-1 level > 2 Standard Deviations (SD) below the mean (*must include chart notes and documentation to confirm diagnosis and lab results with reference ranges*); AND
- 5. Member must have a pretreatment height (*must include growth charts*) of > 2 SD below the mean for age and gender; AND
- If member is age 12 or older, radiographic evidence the member's epiphyses are open (*x-ray results must be included*). Comparison of bone age to chronological age should be documented as abnormal by > 2 SD below the mean for chronological age.
- 7. **Dosage allowed:** 0.18 mg/kg/week.

#### *If member meets all the requirements listed above, the medication will be approved for 12 months.* For **reauthorization**:

- 1. Member must be in compliance with all of the initial criteria; AND
- If member is age 12 or older, radiographic evidence the member's epiphyses are open (*x-ray results must be included*). Comparison of bone age to chronological age should be documented as abnormal by > 2 SD below the mean for chronological age; AND
- Member has a growth rate > 2.5 cm/year unless there is a documented reason for lack of efficacy (on treatment < 1 year, off treatment for a reason for a period of time, nearing final adult height, late stages of puberty).

# *If member meets all the reauthorization requirements above, the medication will be approved for an additional 12 months.*

CareSource considers Saizen (somatropin) not medically necessary for the treatment of the following disease states based on a lack of robust clinical controlled trials showing superior efficacy compared to currently available treatments:

- Constitutional growth delay
- Corticosteroid-induced growth failure
- Cystic fibrosis
- Idiopathic, or non-growth hormone dependent, short stature
- Juvenile idiopathic, or chronic, arthritis
- Noonan Syndrome
- Obesity



- Partial growth hormone deficiency
- Pediatric growth failure due to chronic kidney disease
- Prader-Willi Syndrome
- SHOX deficiency
- Small for Gestational Age
- Turner Syndrome
- Wound healing in burns patients

DATE	ACTION/DESCRIPTION	
10/25/2018	New policy for Saizen created.	

References:

- 1. Saizen [prescribing information]. Rockland, MD: EMD Serono, Inc.; Revised December 2016.
- Nemecheck PM, Polsky B, Gottlieb MS. Treatment Guidelines for HIV-associated wasting. May Clinc Proc. 2000; 27: 386-394.
- Blum WF, Crowe BJ, Quigley CA, et al. Growth hormone in effective in treatment of short stature associated with short stature homeobox-containing gene deficiency: two-year results of a randomized, controlled, multicenter trial. J Clin Endocinol Metab. 2007; 92: 219-228.
- 4. Blum WF, Ross JL, Zimmermann Ag, et al. Growth hormone treatment to final height produces similar height gains in patients with SHOX deficiency and Tuner syndrome: results of a multicenter trial. J Clin Endocrinol Metab. 2013; 98 (8): 1383-1392.
- 5. Kirk J, Betts P, Butler G, et al. Short stature in Noonan syndrome: response to growth hormone therapy. Arch Dis Child. 2001; 84(5): 440-443.
- 6. Raynal P. Growth hormone and noonan syndrome: update in dysfunctional signaling aspects and in therapy for short stature.
- 7. Mahan JD, Warady BA. Assessment and treatment of short stature in pediatric patients with chronic kidney disease: a consensus statement. Pediatr Nephrol. 2006; 21(7): 917-930.
- 8. Romano AA, Allanson JE, Dahlgren J, et al. Noonan syndrome: clinical features, diagnosis, and management guidelines. Pediatrics 2010;126(4): 746-759
- 9. Clayton PE, Cianfarani S, Czernichow P, et al. Management of the Child Born Small for Gestational Age Through to Adulthood: A Consensus Statement of the International Societies of Pediatric Endocrinology and the Growth Hormone Research Society, J Clin Endrocrinol Metab. 2007; 92(3): 804-810.
- 10. Baxter L, Bryant J, Cave CB, Milne R. Recombinant growth hormone for children and adolescents with Turner syndrome.

Effective date: 02/01/2019 Revised date: 10/25/2018



- 1) Acquired structural abnormalities:
  - CNS tumor or neoplasm (craniopharyngioma, glioma, pituitary adenoma, etc.)
  - Cysts (Rathke cleft cyst or arachnoid cleft cyst)
  - Surgery
  - Radiation
  - Chemotherapy
  - CNS infection
  - CNS infarction (e.g., Sheehan's syndrome)
  - Inflammatory lesions (e.g., autoimmune hypohysitis)
  - Infiltrative lesions (e.g., sarcoidosis, histiocytosis)
  - Head trauma or traumatic brain injury
  - Aneurysmal subarachnoid hemorrhage
  - Panhypopituitarism
- 2) Congenital abnormalities:
  - Known genetic mutations in growth-hormone releasing hormone (GHRH) receptor, GH gene, GH receptor or pituitary transcription factors
  - Optic nerve hypoplasia/septo-optic dysplasia
  - Empty sella syndrome
  - Ectopic posterior pituitary
  - Pituitary aplasia/hypoplasia
  - Pituitary stalk defect
  - Anencephaly or prosencephaly
  - Other mid-line defects
  - Vascular malformations
- 3) Pituitary hormones, other than growth hormone (GH):
  - Adrenocorticotropic hormone (ACTH)
  - Antidiuretic hormone (ADH)
  - Follicle stimulating hormone (FSH)
  - Luteinizing hormone (LH)
  - Oxytocin
  - Prolactin
  - Thyroid stimulating hormone (TSH)