

# PHARMACY POLICY STATEMENT

## Marketplace

<b>DRUG NAME</b>	<b>Sohonos (palovarotene)</b>
BENEFIT TYPE	Pharmacy
STATUS	Prior Authorization Required

Sohonos, initially approved by the FDA in 2023, is a retinoid indicated for reduction in the volume of new heterotopic ossification in adults and children aged 8 years and older for females and 10 years and older for males with fibrodysplasia ossificans progressiva (FOP). Through binding to RAR, Sohonos decreases the BMP/ALK2 downstream signaling pathway by inhibiting the phosphorylation of SMAD1/5/8, which reduces ALK2/SMAD-dependent chondrogenesis and osteocyte differentiation resulting in reduced endochondral bone formation.

FOP is an ultra-rare condition that causes abnormal bone growth in areas outside of the skeleton such as ligaments, tendons, and muscles. The disease progresses with flare-up episodes that lead to rapid heterotopic ossification (HO), severely restricting mobility and function as well as quality of life.

Approval was based on the phase 3 MOVE trial which did not meet the primary endpoint of annualized volume of new HO measured by low-dose whole-body computed tomography (WBCT). However, a post hoc 18-month interim analysis showed that Sohonos reduced annualized HO volume by 54% compared with standard of care.

Sohonos (palovarotene) will be considered for coverage when the following criteria are met:

### **Fibrodysplasia Ossificans Progressiva (FOP)**

For **initial** authorization:

1. If member is female, member is at least 8 years of age; OR

***If all the above requirements are met, the medication will be approved for 12 months.***

For **reauthorization**:

1. Chart notes have been provided showing improvement of signs and symptoms of disease (such as reduced volume of new heterotopic ossifications, decreased flare ups, decreased pain or increased mobility); AND
2. If member has not reached skeletal maturity or final adult height, chart notes must include radiological evidence of appropriate bone age (x-ray results must be included) and linear growth.

***If all the above requirements are met, the medication will be approved for an additional 12 months.***

**CareSource considers Sohonos (palovarotene) not medically necessary for the treatment of conditions that are not listed in this document. For any other indication, please refer to the Off-Label policy.**

DATE	ACTION/DESCRIPTION
11/9/2023	New policy for Sohonos created.

References:

1. Sohonos [prescribing information]. Cambridge, MA: Ipsen Biopharmaceuticals, Inc.; 2023.
2. Pignolo RJ, Hsiao EC, Al Mukaddam M, et al. Reduction of New Heterotopic Ossification (HO) in the Open-Label, Phase 3 MOVE Trial of Palovarotene for Fibrodysplasia Ossificans Progressiva (FOP). *J Bone Miner Res.* 2023;38(3):381-394. doi:10.1002/jbmr.4762.
3. Kaplan FS, et al. TmnBT/F9q0.000009s18(p)-58(l)-8(a)13 0 1 72.1 4q7( )-29(e)18(r43.55 430.95 12.8 reWs)-37(:) Tf