[You may adapt this sample Letter of Medical Necessity to reflect specific criteria, including your clinical judgment and accurate patient experience.]

Prescriber Name: *Insert prescriber’s name*

Prescriber Title: *Insert prescriber’s title*

Prescriber Contact Information: *Insert provider’s contact information*

Date: *Insert date*

Insurance Company Name: *Insert insurance company name*

Insurance Company Address: *Insert insurance company address*

Subject: Letter of Medical Necessity for parathyroid hormone therapy for *Insert patient’s name*

To Whom It May Concern:

I am writing to request approval for YORVIPATH® (palopegteriparatide), a parathyroid hormone (PTH) therapy,   
for my patient *Insert patient’s name*, who is suffering from hypoparathyroidism. This therapy is crucial for managing their symptoms. Given the refractory nature of *Insert patient’s name*’s hypocalcemia and the severe impact on   
their health, it is medically necessary to *re-initiate / continue* PTH therapy. This treatment significantly reduces   
their symptoms and prevents further complications associated with chronic hypocalcemia.

**The burden of hypoparathyroidism negatively impacts health-related quality of life (QoL), irrespective   
of serum calcium level, as well as physical functioning and psychological well-being.1,2**

PTH therapy is a critical and necessary treatment for *Insert patient’s name* to manage their condition effectively. Unlike conventional therapy that merely aims to increase calcium absorption, PTH therapy directly addresses the underlying deficiency.

**The 2022 Guidelines from the Second International Workshop state to consider the use of PTH when conventional therapy is deemed unsatisfactory. *Insert patient’s name* meets the following criteria that the guidelines use to define inadequate control with conventional therapy and justify the use of PTH therapy**3**:**

* **Symptomatic hypocalcemia**
* **Hyperphosphatemia**
* **Renal insufficiency**
* **Hypercalciuria**
* **Poor QoL**

**The guidelines also note that patients with chronic hypoparathyroidism with poor compliance, malabsorption, or who are intolerant of large doses of calcium and active vitamin D may benefit from   
PTH therapy. Patients requiring high doses of conventional therapy (ie, calcium > 2 g/day or active   
vitamin D > 2 mcg/day) may also benefit from PTH therapy.3**

Clinical studies and guidelines support the efficacy of PTH therapy in patients with chronic hypoparathyroidism who do not respond adequately to conventional therapy. PTH therapy has been shown to maintain stable calcium levels, reduce the severity and frequency of symptoms, and enhance overall well-being.

**YORVIPATH is a prodrug of PTH(1-34) designed to deliver continuous exposure to active PTH over   
24 hours.4 In the phase 3 clinical trial, 68.9% (42 out of 61) of YORVIPATH-treated patients met the efficacy assessment at week 26 compared with 4.8% (1 out of 21) of patients in the placebo group (treatment difference, 64.2%; 95% confidence interval, 49.5% to 78.8%). The efficacy assessment of the phase 3 clinical trial was the proportion of patients who achieved all of the following at week 26: albumin-corrected serum calcium levels in the normal range (8.3 to 10.6 mg/dL); independence from conventional therapy (defined as requiring no active vitamin D and ≤ 600 mg/day of calcium supplementation, including no use   
of as-needed [PRN] doses) since week 22; no increase in study drug dose since week 22; no missing active vitamin D and calcium data since week 22; and study drug dose was ≤ 30 mcg/day during the entire 26-week treatment period.4**

Enclosed, please find supporting medical documentation, including recent laboratory results, a detailed medical history, and clinical guidelines supporting the use of PTH therapy in similar cases.

I urge you to consider this request favorably and approve the coverage for PTH therapy for *Insert patient’s name*. Should you require any additional information, please do not hesitate to contact me directly at *Insert prescriber’s phone number* or *Insert prescriber’s email address*.

Thank you for your attention to this matter.

Sincerely,

*Insert your name*

*Insert your title*

*Insert your medical institution or practice name*

*Insert your contact information*

Attachments:

1. Patient medical records
2. Recent laboratory results
3. Relevant clinical guidelines and supporting literature

**Please see the diagnosis criteria and attached clinicals:**

**ICD-10 code: *YORVIPATH is indicated for [E20.9, E89.2]***

**PTH level: *Result; Date***

**24-hour urine calcium: *Result; Date***

**Serum calcium level: *Result; Date***

**Serum 25(OH) vitamin D: *Result; Date***

**Daily calcium supplement dose: *mg elemental calcium/day***

**Daily active vitamin D (calcitriol) dose: *mcg/day; Date***

**Date of surgery: *Date***

**Clinical risks persisting under current conventional therapy:**

* **Renal complications** 
  + Patients with chronic hypoparathyroidism who receive long-term treatment with conventional therapy   
    (oral calcium and active vitamin D) have an increased risk of renal complications compared to the general population5-7
  + Up to 41% (44 out of 107) of patients with chronic hypoparathyroidism treated with conventional therapy have reported chronic kidney disease (CKD), and 36% (92 out of 259) have reported nephrolithiasis5-7
* **Skeletal** 
  + Hypoparathyroidism is associated with reduced bone turnover and high bone mineral density8,9
* **Cardiovascular** 
  + Longer duration of hypoparathyroidism, lower time-weighted serum ionized calcium, and the occurrence of 4 or more hypercalcemic episodes all significantly increased the risk of developing cardiovascular disease in a case-control study of 431 patients with hypoparathyroidism treated with conventional therapy10
* **Neuropsychiatric**
  + In interviews conducted during the development of the Hypoparathyroidism Patient Experience Scale (HPES), a majority of patients reported reduced functioning and well-being despite treatment with conventional therapy, including having anxiety (81% or 34 out of 42), feeling sad or depressed (62% or   
    26 out of 42), and feeling irritable or short-tempered (43% or 18 out of 42).2 The HPES was developed through interviews of 5 clinical experts and 42 adult participants with hypoparathyroidism to evaluate the symptoms (physical and cognitive) and impact (characterized as physical functioning, psychological   
    well-being, daily life, and social life and relationships) associated with hypoparathyroidism2,11

**References: 1.** Kontogeorgos G, Mamasoula Z, Krantz E, Trimpou P, Landin-Wilhelmsen K, Laine CM. Low health-related quality of life in hypoparathyroidism and need for PTH analog. *Endocr Connect.* 2022;11(1):e210379. doi:10.1530/EC-21-0379 **2.** Brod M, Waldman LT, Smith A, Karpf D. Living with hypoparathyroidism: development of the Hypoparathyroidism Patient Experience Scale-Impact (HPES-Impact). *Qual Life Res.* 2021;30(1):277-291. doi:10.1007/s11136-020-02607-1   
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**5.** Gosmanova EO, Houillier P, Rejnmark L, Marelli C, Bilezikian JP. Renal complications in patients with chronic hypoparathyroidism on conventional therapy:   
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of hypoparathyroidism: etiologies and clinical features. *J Clin Endocrinol Metab.* 2016;101(6):2300-2312. doi:10.1210/jc.2015-3909 **9.** Rubin MR, Dempster DW, Zhou H, et al. Dynamic and structural properties of the skeleton in hypoparathyroidism. *J Bone Miner Res.* 2008;23(12):2018-2024. doi:10.1359/JBMR.080803   
**10.** Underbjerg L, Sikjaer T, Rejnmark L. Long-term complications in patients with hypoparathyroidism evaluated by biochemical findings: a case-control study.   
*J Bone Miner Res.* 2018;33(5):822-831. doi:10.1002/jbmr.3368 **11.** Brod M, Waldman LT, Smith A, Karpf D. Assessing the patient experience of hypoparathyroidism symptoms: development of the Hypoparathyroidism Patient Experience Scale-Symptom (HPES-Symptom). *Patient.* 2020;13(2):151-162. doi:10.1007/s40271-  
019-00388-5

**Important Safety Information**

**INDICATION AND LIMITATIONS OF USE**

YORVIPATH (palopegteriparatide) is indicated for the treatment of hypoparathyroidism in adults.

* YORVIPATH was not studied for acute post-surgical hypoparathyroidism.
* YORVIPATH’s titration scheme was only evaluated in adults who first achieved an albumin-corrected serum calcium of at least 7.8 mg/dL using calcium and active vitamin D treatment.

**CONTRAINDICATIONS**

YORVIPATH is contraindicated in patients with severe hypersensitivity to palopegteriparatide or to any of its excipients. Hypersensitivity reactions, including anaphylaxis, angioedema, and urticaria, have been observed with parathyroid hormone (PTH) analogs.

**WARNINGS AND PRECAUTIONS**

**Risk of Unintended Changes in Serum Calcium Levels Related to Number of Daily Injections**

Use only one YORVIPATH injection to achieve the recommended once daily dosage. Using two YORVIPATH injections to achieve the recommended once daily dosage increases the variability of the total delivered dose, which can cause unintended changes in serum calcium levels, including hypercalcemia and hypocalcemia.

**Serious Hypercalcemia**

Serious events of hypercalcemia requiring hospitalization have been reported with YORVIPATH. The risk is   
highest when starting or increasing the dose of YORVIPATH but may occur at any time. Measure serum calcium   
7 to 10 days after any dose change or if there are signs or symptoms of hypercalcemia, and at a minimum of every 4 to 6 weeks once the maintenance dose is achieved. Treat hypercalcemia if needed. If albumin-corrected serum calcium is greater than 12 mg/dL, withhold YORVIPATH for at least 2-3 days. For less serious hypercalcemia, adjust the dose of YORVIPATH, active vitamin D, and/or calcium supplements.

**Serious Hypocalcemia**

Serious events of hypocalcemia have been observed with PTH products, including YORVIPATH. The risk is highest when YORVIPATH is abruptly discontinued, but may occur at any time, even in patients who have been on stable doses of YORVIPATH. Measure serum calcium 7 to 10 days after any dose change or if there are signs or symptoms of hypocalcemia, and at a minimum of every 4 to 6 weeks once the maintenance dosage is achieved. Treat hypocalcemia if needed, and adjust the dose of YORVIPATH, active vitamin D, and/or calcium supplements if hypocalcemia occurs.

**Potential Risk of Osteosarcoma**

YORVIPATH is a PTH analog. An increased incidence of osteosarcoma (a malignant bone tumor) has been reported in male and female rats treated with PTH analogs, including teriparatide. Osteosarcoma occurrence in rats is dependent on teriparatide or PTH dose and treatment duration. Osteosarcoma has been reported in patients treated with teriparatide in the postmarketing setting; however, an increased risk of osteosarcoma has not been observed in observational studies in humans. There are limited data assessing the risk of osteosarcoma beyond   
2 years of teriparatide use.

YORVIPATH is not recommended in patients who are at increased risk of osteosarcoma, such as patients with:

* Open epiphyses. YORVIPATH is not approved in pediatric patients.
* Metabolic bone diseases other than hypoparathyroidism, including Paget’s disease of bone.
* Unexplained elevations of alkaline phosphatase.
* Bone metastases or a history of skeletal malignancies.
* History of external beam or implant radiation therapy involving the skeleton.
* Hereditary disorders predisposing to osteosarcoma.

Instruct patients to promptly report clinical symptoms (e.g., persistent localized pain) and signs   
(e.g., soft tissue mass tender to palpation) that could be consistent with osteosarcoma.

**Orthostatic Hypotension**

Orthostatic hypotension has been reported with YORVIPATH. Associated signs and symptoms may include decreased blood pressure, dizziness (including postural dizziness), palpitations, tachycardia, presyncope, or syncope. Such symptoms can be managed by dosing at bedtime, while reclining. YORVIPATH should be administered initially when the patient can sit or lie down due to the potential of orthostatic hypotension.

**Risk of Digoxin Toxicity with Concomitant Use of Digitalis Compounds**

YORVIPATH increases serum calcium, and therefore, concomitant use with digoxin (which has a narrow therapeutic index) may predispose patients to digitalis toxicity if hypercalcemia develops. Digoxin efficacy may be reduced if hypocalcemia is present. When YORVIPATH is used concomitantly with digoxin, measure serum calcium and digoxin levels routinely, and monitor for signs and symptoms of digoxin toxicity. Refer to the digoxin prescribing information for dose adjustments, if needed.

**ADVERSE REACTIONS**

The most common adverse reactions (≥ 5%) in patients treated with YORVIPATH were injection site   
reactions (39%), vasodilatory signs and symptoms (28%), headache (21%), diarrhea (10%), back pain (8%), hypercalcemia (8%) and oropharyngeal pain (7%).

**DRUG INTERACTIONS**

**Drugs Affected by Serum Calcium**

Digoxin: YORVIPATH increases serum calcium, therefore, concomitant use with digoxin (which has a narrow therapeutic index) may predispose patients to digitalis toxicity if hypercalcemia develops. Digoxin efficacy may be reduced if hypocalcemia is present. When YORVIPATH is used concomitantly with digoxin, measure serum calcium and digoxin levels, and monitor for signs and symptoms of digoxin toxicity. Adjustment of the digoxin and/or YORVIPATH dose may be needed.

**Drugs Known to Affect Serum Calcium**

Drugs that affect serum calcium may alter the therapeutic response to YORVIPATH. Measure serum calcium more frequently when YORVIPATH is used concomitantly with these drugs, particularly after these drugs are initiated, discontinued, or dose adjusted.

**USE IN SPECIFIC POPULATIONS**

**Pregnancy**

Available data from reports of pregnancies in the clinical trials from drug development are insufficient to identify   
a drug-associated risk of major birth defects, miscarriage, or other adverse maternal or fetal outcomes. If YORVIPATH is administered during pregnancy, or if a patient becomes pregnant while receiving YORVIPATH, healthcare providers should report YORVIPATH exposure by calling 1-844-442-7236.

**Lactation**

Monitor infants breastfed by females treated with YORVIPATH for symptoms of hypercalcemia or hypocalcemia. Consider monitoring serum calcium in the breastfed infant.

You are encouraged to report side effects to FDA at (800) FDA-1088 or www.fda.gov/medwatch.   
You may also report side effects to Ascendis Pharma at 1-844-442-7236.

**Please see Important Safety Information on pages 4 and 5 and full Prescribing Information for   
YORVIPATH at YorvipathHCP.com.**