HEALTH PARTNERS PLANS 2024 PRIOR AUTHORIZATION REQUEST FORM

A part of Jefferson Health Plans

Bone Density Regulators

Phone: 215-991-4300

Fax back to: 866-240-3712

Health Partners Plans manages the pharmacy drug benefit for your patient. Certain requests for coverage require review with the prescribing physician. Please answer the following questions and fax this form to the number listed above.

PLEASE NOTE: Any information (patient, prescriber, drug, labs) left blank, illegible, or not attached WILL DELAY the review process.

Patient Name:	Prescriber Name:	
HPP HPP Member Number:	Fax:	Phone:
Date of Birth:	Office Contact:	
Patient Primary Phone:	NPI:	PA PROMISe ID:
Address:	Address:	
City, State ZIP:	City, State ZIP:	
Line of Business: Medicaid CHIP	Specialty Pharmacy (if applicable):	
Drug Name:	Strength:	
Quantity:	Refills:	
Directions:		
Diagnosis Code: Diagnosis:		
HPP's maximum approval time is 12 mo	onths but may be less dependin	g on the drug.
Please attach any pertinent medical history including lab Please answer the fol	s and information for this me lowing questions and sign.	mber that may support approval.
Q1. Is the requested medication prescribed for the treatment of a diagnosis that is indicated in the U.S. Food and Drug Administration (FDA)-approved package labeling OR a medically accepted indication?		
□ Yes	□ Yes □ No	
Q2. Is the prescribed dose and duration of therapy for the requested medication consistent with FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature?		
□ Yes □ No		
Q3. Does the patient have a contraindication to the requested medication?		
□ Yes	□ No	
Q4. Is the request for an osteoporosis related condition?		
□ Yes	□ No	

Q5. Was the patient evaluated for secondary causes of osteoporosis including the following: complete blood count (CBC), vitamin D, ionized calcium, phosphorus, albumin, total protein,

HEALTH PARTNERS PLANS 2024 PRIOR AUTHORIZATION REQUEST FORM

A part of Jefferson Health Plans

Bone Density Regulators

Phone: 215-991-4300

Fax back to: 866-240-3712

Health Partners Plans manages the pharmacy drug benefit for your patient.	Certain requests for coverage require review with the prescribing physician.
Please answer the following questions and fax this form to the number lister	d above.

PLEASE NOTE: Any information (patient, prescriber, drug, labs) left blank, illegible, or not attached WILL DELAY the review process.

creatinine, liver enzymes (specifically alkaline phosphatase), intact parathyroid hormone (PTH), thyroid stimulating hormone (TSH), urinary calcium excretion, and testosterone (if a male)? PYes No Q6. Is the request for an anabolic agent? No Q7. Does the patient have a T score of negative three and five tenths (-3.5) or below? Yes Q8. Does the patient have a T score of negative two and five tenths (-2.5) or below? No Q8. Does the patient have a T score of negative two and five tenths (-2.5) or below? No Q9. Does the patient have a history of fragility fracture? No Q10. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Yes No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? Yes No Q12. Does the patient have a history of Skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	Patient Name:	Prescriber Name:
Q6. Is the request for an anabolic agent? Pres No Q7. Does the patient have a T score of negative three and five tenths (-3.5) or below? Pres No Q8. Does the patient have a T score of negative two and five tenths (-2.5) or below? Pres No Q9. Does the patient have a T score of negative two and five tenths (-2.5) or below? Pres No Q9. Does the patient have a history of fragility fracture? Pres No Q10. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Pres No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? Pres No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?		
Yes No Q7. Does the patient have a T score of negative three and five tenths (-3.5) or below? Yes No Q8. Does the patient have a T score of negative two and five tenths (-2.5) or below? Yes No Q9. Does the patient have a history of fragility fracture? Yes No Q9. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Yes No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? Yes No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	□ Yes	□ No
Q7. Does the patient have a T score of negative three and five tenths (-3.5) or below? Pres No Q8. Does the patient have a T score of negative two and five tenths (-2.5) or below? Pres No Q9. Does the patient have a history of fragility fracture? Pres No Q10. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Pres No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? Pres No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	Q6. Is the request for an anabolic agent?	
□ Yes □ No Q8. Does the patient have a T score of negative two and five tenths (-2.5) or below? □ Yes □ No Q9. Does the patient have a history of fragility fracture? □ Yes □ No Q10. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? □ Yes □ No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? □ Yes □ No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	☐ Yes	□ No
Q8. Does the patient have a T score of negative two and five tenths (-2.5) or below? Yes No Q9. Does the patient have a history of fragility fracture? No Yes No Q10. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Yes No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? Yes No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	Q7. Does the patient have a T score of negative	three and five tenths (-3.5) or below?
□ Yes □ No Q9. Does the patient have a history of fragility fracture? □ Yes □ Yes □ No Q10. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? □ Yes □ No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? □ Yes □ No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	□ Yes	□ No
Q9. Does the patient have a history of fragility fracture? Pres No Q10. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Pres No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? Yes No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	Q8. Does the patient have a T score of negative two and five tenths (-2.5) or below?	
□ Yes □ No Q10. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? □ Yes □ No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? □ Yes □ No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	□ Yes	□ No
Q10. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? □ Yes □ No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? □ Yes □ No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	Q9. Does the patient have a history of fragility fracture?	
or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Yes No Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? Yes No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	□ Yes	□ No
Q11. Has the patient received a cumulative treatment duration that exceeds recommendations in the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? Yes No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	or fragility fracture after two (2) or more years despite treatment with a bisphosphonate),	
the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature? Yes No Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	□ Yes	□ No
Q12. Does the patient have a history of ANY of the following: For Forteo and Tymlos: A) Paget's disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	the FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical	
disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving the skeleton; for Evenity: A) myocardial infarction?	□ Yes	□ No
□ Yes □ No	disease, B) bone metastases, C) a history of skeletal malignancies, D) metabolic bone disease other than osteoporosis, E) hypercalcemic disorders, F) unexplained elevations of alkaline phosphatase, G) open epiphyses, or H) prior external beam or implant radiation therapy involving	
		□ No

HEALTH PARTNERS PLANS 2024 PRIOR AUTHORIZATION REQUEST FORM

A part of Jefferson Health Plans

Bone Density Regulators

Phone: 215-991-4300

Fax back to: 866-240-3712

Health Partners Plans manages the pharmacy drug benefit for your patient.	Certain requests for coverage require review with the prescribing physician.
Please answer the following questions and fax this form to the number liste	d above.

PLEASE NOTE: Any information (patient, prescriber, drug, labs) left blank, illegible, or not attached WILL DELAY the review process.

Patient Name:	Prescriber Name:
Q13. Is the requested medication Evenity (romosozumab) or Tymlos (abaloparatide)?	
□ Yes	□ No
Q14. Is the requested medication teriparatide?	
□ Yes	□ No
Q15. Is the requested medication Forteo?	
□ Yes	□ No
Q16. Does the patient have a contraindication or intolerance to that would not be expected to occur with Forteo?	
□ Yes	□ No
Q17. Is the requested medication Evista (raloxifene)?	
□ Yes	□ No
Q18. Does not have a history of venous thromboembolic event or breast cancer? Yes	
□ Yes	□ No
Q19. Is the patient a women with a risk factor for stroke (such as prior stroke or transient ischemic attack (TIA), atrial fibrillation, hypertension, or cigarette smoking)?	
□ Yes	□ No
Q20. Has the increased risk of death due to stroke been discussed with the patient and documented by the prescriber?	
□ Yes	□ No
Q21. Is the patient a postmenopausal woman at high risk of fracture (i.e., T-score between negative 1.0 and negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 3 percent or a 10-year probability of a major osteoporosis-related fracture greater than or equal to 20 percent	
	et is to well a maintenant. This information is informated and a factor for the same of the individual or

HEALTH PARTNERS PLANS 2024 PRIOR AUTHORIZATION REQUEST FORM

A part of Jefferson Health Plans

Bone Density Regulators

Phone: 215-991-4300

Fax back to: 866-240-3712

Health Partners Plans manages the pharmacy drug benefit for your patient. Certain requests for coverage require review with the prescribing physician. Please answer the following questions and fax this form to the number listed above.

PLEASE NOTE: Any information (patient, prescriber, drug, labs) left blank, illegible, or not attached WILL DELAY the review process.

based on the US-adapted World Health Organization (WHQ) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)? Yes No Q22. Is the patient at high risk for invasive breast cancer as defined by one of the following: A) prior biopsy with lobular carcinoma in situ (LCIS) or atypical hyperplasia, B) one or more first degree relatives with breast cancer, or C) A 5-year predicted risk of breast cancer greater than or equal to 1.66 percent (based on the modified Gail model)? Yes No Q23. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Yes No Q24. Is the requested medication Xgeva (denosumab)? No Yes No Q25. Does the patient have a high risk of fracture (i.e., T-score between negative 1.0 and negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 3 percent or a 10-year probability of a major osteoporosis-related fracture greater than or equal to 20 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)? Yes No <	Patient Name:	Prescriber Name:
Q22. Is the patient at high risk for invasive breast cancer as defined by one of the following: A) prior biopsy with lobular carcinoma in situ (LCIS) or atypical hyperplasia, B) one or more first degree relatives with breast cancer, or C) A 5-year predicted risk of breast cancer greater than or equal to 1.66 percent (based on the modified Gail model)? □ Yes □ No Q23. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? □ Yes □ No Q24. Is the requested medication Xgeva (denosumab)? □ Yes □ Yes □ No Q25. Does the patient have a high risk of fracture (i.e., T-score between negative 1.0 and negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 and a history of fragility fracture greater than or equal to 3 percent or a 10-year probability of a hip fracture greater than or equal to 2 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)? □ Yes □ No Q26. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to the preferred bone density regulators approved or medically accepted for the patient's diagnosis?	below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip	
prior biopsy with lobular carcinoma in situ (LCIS) or atypical hyperplasia, B) one or more first degree relatives with breast cancer, or C) A 5-year predicted risk of breast cancer greater than or equal to 1.66 percent (based on the modified Gail model)? Yes No Q23. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Yes No Q24. Is the requested medication Xgeva (denosumab)? No Q25. Does the patient have a high risk of fracture (i.e., T-score between negative 1.0 and negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 20 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)? Yes No	□ Yes	□ No
Q23. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? □ Yes □ No Q24. Is the requested medication Xgeva (denosumab)? □ No Q25. Does the patient have a high risk of fracture (i.e., T-score between negative 1.0 and negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 3 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)? □ Yes □ No Q26. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to the preferred bone density regulators approved or medically accepted for the patient's diagnosis?	prior biopsy with lobular carcinoma in situ (LCIS) or atypical hyperplasia, B) one or more first degree relatives with breast cancer, or C) A 5-year predicted risk of breast cancer greater than or	
or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to bisphosphonates? Yes Q24. Is the requested medication Xgeva (denosumab)? Yes Q25. Does the patient have a high risk of fracture (i.e., T-score between negative 1.0 and negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 3 percent or a 10-year probability of a major osteoporosis-related fracture greater than or equal to 20 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)? Yes Q26. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to the preferred bone density regulators approved or medically accepted for the patient's diagnosis?	□ Yes	□ No
Q24. Is the requested medication Xgeva (denosumab)? Yes No Q25. Does the patient have a high risk of fracture (i.e., T-score between negative 1.0 and negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 3 percent or a 10-year probability of a major osteoporosis-related fracture greater than or equal to 20 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)? Yes No Q26. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to the preferred bone density regulators approved or medically accepted for the patient's diagnosis?	or fragility fracture after two (2) or more years despite treatment with a bisphosphonate),	
□ Yes □ No Q25. Does the patient have a high risk of fracture (i.e., T-score between negative 1.0 and negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 3 percent or a 10-year probability of a major osteoporosis-related fracture greater than or equal to 20 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)? □ Yes □ No Q26. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to the preferred bone density regulators approved or medically accepted for the patient's diagnosis?	□ Yes	□ No
Q25. Does the patient have a high risk of fracture (i.e., T-score between negative 1.0 and negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 3 percent or a 10-year probability of a major osteoporosis-related fracture greater than or equal to 20 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)?	Q24. Is the requested medication Xgeva (denose	umab)?
negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 3 percent or a 10-year probability of a major osteoporosis-related fracture greater than or equal to 20 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture, regardless of bone density)? Q26. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to the preferred bone density regulators approved or medically accepted for the patient's diagnosis?	□ Yes	□ No
Q26. Does the patient have a history of therapeutic failure (i.e., documented continued bone loss or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to the preferred bone density regulators approved or medically accepted for the patient's diagnosis?	negative 2.5 and a history of fragility fracture of the proximal humerus, pelvis, or distal forearm; T-score between negative 1.0 and negative 2.5 at the femoral neck, total hip, or lumbar spine and a 10-year probability of a hip fracture greater than or equal to 3 percent or a 10-year probability of a major osteoporosis-related fracture greater than or equal to 20 percent based on the USadapted World Health Organization (WHO) algorithm; T-score negative 2.5 or below at the femoral neck, total hip, or lumbar spine; OR history of low-trauma spine or hip fracture,	
or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to the preferred bone density regulators approved or medically accepted for the patient's diagnosis?		□ No
□ Yes	or fragility fracture after two (2) or more years despite treatment with a bisphosphonate), intolerance, or contraindication to the preferred bone density regulators approved or medically	
	□ Yes	□ No

HEALTH PARTNERS PLANS 2024 PRIOR AUTHORIZATION REQUEST FORM

A part of Jefferson Health Plans

Bone Density Regulators

Phone: 215-991-4300

Fax back to: 866-240-3712

Health Partners Plans manages the pharmacy drug benefit for your patient. Certain requests for coverage require review with the prescribing physician. Please answer the following questions and fax this form to the number listed above.

PLEASE NOTE: Any information (patient, prescriber, drug, labs) left blank, illegible, or not attached WILL DELAY the review process.

Patient Name:	Prescriber Name:
Q27. Is the request for a parenteral bisphosphonate?	
□ Yes	□ No
Q28. Does the patient have a contraindication or intolerance to oral bisphosphonates?	
□ Yes	□ No
Q29. For renewals: Has the patient's condition stabilized or is the patient benefitting from the requested drug?	
□ Yes	□ No
Q30. Additional Information:	

Prescriber Signature

Date

Updated for 2024