

New Clinically-Proven Treatment for Osteopenia

The first and only non-pharmacological prescription treatment for your patients with low bone density that has clinically-proven effectiveness.



FDA-Cleared Precision Vibration Therapy

- **Clinically-Proven Efficacy:** Clinical trial demonstrated significant reduction in the loss of bone density and strength in postmenopausal women with osteopenia.
- **High Patient Compliance:** 80% adherence to the treatment regimen.
- **Targeted & Calibrated Vibration:** Embedded sensors provide targeted, calibrated vibration therapy to the spine and hips.

Preserve Bone Density and Strength

Clinical Trial in Postmenopausal Women with Osteopenia (n=126)¹

83% reduction in bone strength loss

85% reduction in bone density loss

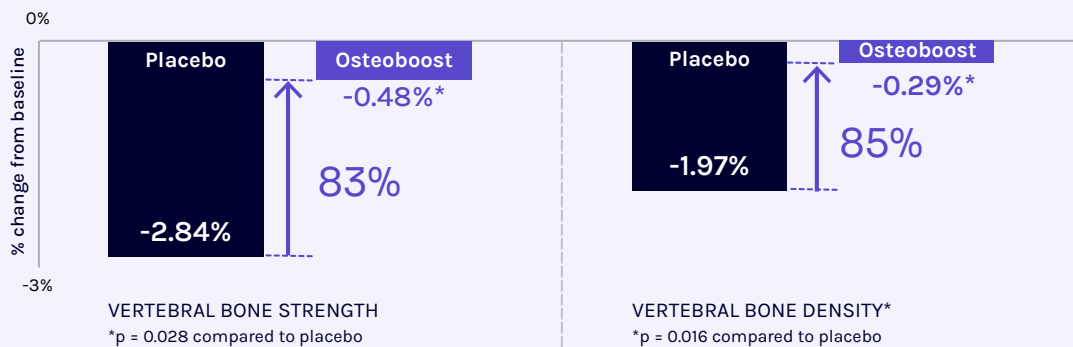


Figure shows data from participants who averaged 3 or more treatments per week over 12 months (n=73 participants)

1. Bilek LD, Flores LE, Waltman N, et al. Benefits of Targeted Vibration for Bone Strength and Bone Density in Postmenopausal Women with Osteopenia: A Randomized, Sham-Controlled Trial. JBMR Plus. 2024;ziae104. doi:10.1093/jbmrpl/ziae104



Scan here for the JBMR Publication



For more on Osteoboost scan here or go to osteoboost.com

Targeted, Precise Therapy

- **Targeted vibration** directly to the spine & hips provides mechanical stimulation that mimics the mechanism and effect of high-impact exercise
- **Reduces the loss of the bone density and bone strength**
- **Embedded sensors** ensure therapeutic vibration dose during every treatment session



Increased Usage Leads to Even Greater Benefit

More frequent use of Osteoboost is correlated with greater benefit, with 31% of women even experiencing an **increase in bone strength**.

Preventing Osteoporosis

The clinical trial treated patients for 12 months. Figure (left) illustrates a multiyear projection based on the study results at 12 months.

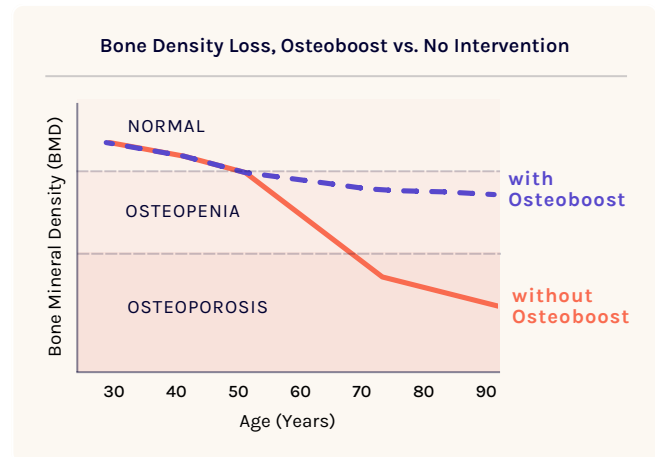
How to Use Osteoboost

Osteoboost is comfortable and as easy to use as putting on a belt. Each daily treatment session is **30 minutes**. Osteoboost **can be worn while doing chores or walking**, making it easy to fit into any lifestyle. Osteoboost should be used at least five days per week. For best results, we recommend that Osteoboost be used daily.

Safety & Compliance

In our 12-month clinical trial, there were no device-related serious adverse effects. The most common adverse effects potentially related to the device were back pain, pelvic pain, and leg pain, experienced in both the Active and Placebo groups. These were usually mild and transient.

Overall compliance was 80% and participants found the belt comfortable and easy to use.



Prescribing Osteoboost

Fax: 1-888-870-2808 • Phone: 1-800-748-7001

E-Prescribe: healthwarehouse.com (Florence, KY) • NPI: 1619252160 • NCPDP#: 1832674

Diagnosis ICD-10 Code: ICD M85. 8 (Osteopenia)

For ordering, please direct patients to visit Osteoboost.com

Osteoboost is currently available through self-pay. We are working to secure coverage from major insurers and Medicare.



Scan here for Safety & Indications



For more on Osteoboost scan here or go to osteoboost.com



Patient Information

Name	D.O.B.
Phone	Email
Address	City
State	Zip Code

Prescriber Information

Prescriber Name	NPI
Address	City
State	Zip Code
Office Contact	Email
Phone	Fax

Prescription information: Osteoboost

Check the appropriate ICD-10 Code:

 M85. 8 (Osteopenia) Other**Usage:** 30 minutes per day, daily.Clinical criteria: Before use, refer to the product labeling for complete product instructions for use, warnings, and precautions at osteoboost.com/indications-for-use

Prescriber Authorization

I certify that Osteoboost is reasonable and medically necessary for the treatment of this patient.

Signature	Date
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LBL-09-2003 rev 1.0, Oct 2024

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