Regional improvements in lumbosacropelvic Hounsfield units following teriparatide treatment

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OBJECTIVE Opportunistic Hounsfield unit (HU) determination from CT imaging has been increasingly used to estimate bone mineral density (BMD) in conjunction with assessments from dual energy x-ray absorptiometry (DXA). The authors sought to compare the effect of teriparatide on HUs across different regions in the pelvis, sacrum, and lumbar spine, as a surrogate measure for the effects of teriparatide on lumbosacropelvic instrumentation.

METHODS A single-institution retrospective review of patients who had been treated with at least 6 months of teriparatide was performed. All patients had at least baseline DXA as well as pre- and post-teriparatide CT imaging. HUs were measured in the pedicle, lamina, and vertebral body of the lumbar spine, in the sciatic notch, and at the S1 and S2 levels at three different points (ilium, sacral body, and sacral ala).

RESULTS Forty patients with an average age of 67 years underwent a mean of 20 months of teriparatide therapy. Mean HUs of the lumbar lamina, pedicles, and vertebral body were significantly different from each other before teriparatide treatment: 343 ± 114, 219 ± 89.2, and 111 ± 48.1, respectively (p < 0.001). Mean HUs at the S1 level for the ilium, sacral ala, and sacral body were also significantly different from each other: 124 ± 90.1, −10.7 ± 61.9, and 99.1 ± 72.1, respectively (p < 0.001). The mean HUs at the S2 level for the ilium and sacral body were not significantly different from each other, although the mean HU at the sacral ala (−11.9 ± 52.6) was significantly lower than those at the ilium and sacral body (p = 0.003 and 0.006, respectively). HU improvement occurred in most regions following teriparatide treatment. In the lumbar spine, the mean lamina HU increased from 343 to 400 (p < 0.001), the mean pedicle HU increased from 219 to 242 (p = 0.04), and the mean vertebral body HU increased from 111 to 134 (p < 0.001). There were also significant increases in the S1 sacral body (99.1 to 130, p < 0.05), S1 ilium (124 vs 165, p = 0.01), S1 sacral ala (−10.7 vs 3.68, p = 0.04), and S2 sacral body (168 vs 189, p < 0.05).

CONCLUSIONS There was significant regional variation in lumbar and sacropelvic HUs, with most regions significantly increasing following teriparatide treatment. The sacropelvic area had lower HU values than the lumbar spine, more regional variation, and a higher degree of correlation with BMD as measured on DXA. While teriparatide treatment resulted in HUs > 110 in the majority of the lumbosacral spine, the HUs in the sacral ala remained suggestive of severe osteoporosis, which may limit the effectiveness of fixation in this region.

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Low bone mineral density (BMD) may be present in as many as 75% of patients undergoing spine surgery, with even higher rates in women older than 50 years. Improving a patient’s BMD prior to surgery is recognized as an important factor in preventing postoperative complications such as pseudarthrosis, junctional kyphosis, pelvic insufficiency fractures below a lumbosacral fusion, and instrumentation failure. Dual-energy x-ray absorptiometry (DXA) is the most commonly utilized method for measuring BMD, but the clinical valida-
tion of this technique has almost exclusively been directed at its ability to assess osteoporotic fracture risk, and it is an imperfect tool. Given the prevalence and morbidity of diminished bone quality in patients undergoing spinal fusion surgery and the limited predictive value of BMD for osteoporosis-related complications of spine surgery, recent work has focused on finding new methods or combinations of methods to better assess bone quality in these patients.

Hounsfield units (HUs) are a relative measurement of x-ray attenuation obtained from routine CT images. Such images are obtained in the vast majority of patients undergoing elective spinal fusion surgery, and HUs have been correlated to BMD assessed by DXA. Thus, HU determination, which can be performed by the surgeon on essentially all PACS systems, represents an opportunistic and essentially costless tool for preoperatively assessing bone quality that has the added benefit of being able to measure more specific regions of interest (ROIs; i.e., vertebral body or pedicle isthmus). An HU cutoff of < 110 in the lumbar spine has been reported to be indicative of osteoporosis. Given degenerative changes and deformity in the spine, the lumbar BMD should not be used and thus is not available to assess bone health in many spine patients. Therefore, HU measurements may be a better tool for use in the lumbar spine and may be superior when assessing the risk for vertebral fracture. HUs allow for more focused assessment of the trabecular bone based on a defined ROI and eliminate the cortical bone, sclerotic changes, and posterior element bone that may affect DXA measurements. However, HUs and spine and hip DXA have been used as complementary or standalone methods to evaluate bone quality.

Teriparatide, a parathyroid hormone (PTH) analog (PTH 1–34), is an anabolic medication that promotes differentiation of osteoblasts, resulting in new bone formation, and is currently approved for the treatment of osteoporosis. Many spine surgeons, often working in consort with bone health specialists such as endocrinologists, have used teriparatide in an off-label fashion to mitigate poor bone quality prior to elective spine surgery. Although there is evidence that antiresorptive therapies may be beneficial for improving fusion rates and decreasing subsequent vertebral column fractures, there is limited evidence that teriparatide is superior to bisphosphonates in promoting fusion and increasing fusion rate, while improving bone quality prior to elective spine surgery. Although there is limited evidence that teriparatide is superior to bisphosphonates in promoting fusion and increasing fusion rate, there is an increased risk with sagittal imbalance, a high pelvic incidence, obesity, and osteoporosis. The sacropelvic area is of particular interest in that sacropelvic fixation has been considered essential to optimize construction stability for long-segment constructs involving the lumbar spine. In a randomized trial of patients with symptomatic pelvic insufficiency fractures, Peichl et al. reported that PTH 1–84 significantly reduced pain, improving walking ability and bony healing compared to placebo. Thus, the use of such an anabolic medication may be useful to prevent and treat such complications related to spinal surgery. With the renewed attention on sacroiliac joint fixation and fusion, it is important to better understand the regional variation in bone density in the sacrum and ilium and any potential benefits of teriparatide.

We hypothesized that a minimum of 6 months of teriparatide treatment would result in significant improvement in the regional bone density of the lumbosacral spine and ilium, using HUs as a surrogate for BMD. Therefore, the objective of this study was to compare the effect of teriparatide on HUs across different regions in the lumbar spine and sacropelvic region. Secondly, we correlated HU changes at each spinal level and anatomical bony region with DXA BMD.

Methods

After obtaining institutional review board approval, we performed a retrospective review of records dated 2002–2018 across all campuses at a single multicenter institution. We identified patients who had been prescribed teriparatide, had both pre- and posttreatment DXA, and had pre- and posttreatment abdominal or lumbar spine CT, including the pelvis. Inclusion criteria were treatment with at least 6 months of teriparatide and CT encompassing all spinal ROIs both before and after treatment. The pre- and posttreatment CT scans had been obtained within a 3-year period. Collected data included age, height, weight, sex, BMI, osteoporosis medications, smoking status, metabolic laboratory tests, hemodialysis status, duration of teriparatide treatment, time between CT scans, time between DXA scans, and BMD at various anatomical regions on DXA. We also manually measured HUs in various ROIs on the CT. HUs were measured using ROI circles drawn in accordance with a previously reported methodology. Two independent reviewers performed all HU measurements.

HU Measurements

For each patient, the following measurements were made using axial scans (Fig. 1) for the lumbar vertebrae: laminar cancellous bone (2 sides, 5 levels), pedicles (2 sides, 5 levels), and vertebral body (3 measurements averaged, 5 levels). The following measurements were made at S1–2 using axial scans: ilium (2 sides, 2 levels), sacral ala (2 sides, 2 levels), and sacral body (2 levels; Fig. 2). Finally, coronal CT images were utilized to measure the sciatic notches (2 sides). When measuring the vertebral body, three ROIs (just inferior to the superior endplate, mid, and just superior to the inferior endplate) were averaged. Mea-
measurements of the iliac crest were made via bilateral ROI ellipses within the trabecular bone at the S2 level. In the lumbar spine, the values from each level were combined to determine a mean value for the vertebral body, pedicle, and lamina. We excluded measurements of vertebrae containing instrumentation, adjacent to interbody grafts, or with vertebroplasty cement, as the artifact confounded HU measurement.

Statistical Analysis

Means were compared via two-sample t-tests, paired t-tests, and one-way ANOVA, with statistical significance defined as $p < 0.05$. Pearson correlation coefficients were calculated to assess the correlation between HUs and DXA BMD before and after teriparatide therapy at each region (lumbar spine and sacropelvic area). Two independent reviewers performed HU measurements, and intraclass correlation coefficient calculations were used to determine intrarater reliability. Analyses were performed using statistical software (JMP Pro 14.0.0, SAS Institute Inc.).

Results

Patient Cohort

We identified a total of 5487 patients who had been prescribed teriparatide over the study period, and 403 of these patients had undergone DXA scanning and lumbar CT imaging. Out of those patients, only 52 had additional CT imaging after 6 months of teriparatide treatment, and 40 patients had imaging adequately encompassing the sacrum and pelvis and were included in the present study. There were 6 men and 34 women with a mean age of 67 ± 12 years (mean ± SD). The mean teriparatide treatment duration was 20 ± 6 months (Table 1). The mean BMI was 24 ± 6 kg/m². The mean femoral neck, lumbar spine, and total hip DXA T-scores were −2.6 ± 0.9, −2.4 ± 1.5, and −2.2 ± 1.1, respectively.

HU Regional Variability Prior to Teriparatide Administration

There were significant differences in HU intensity among regions in the lumbar spine with the lumbar vertebral body, lamina, and pedicle having 111 ± 48.1, 343 ± 114, and 219 ± 89.2 HU, respectively ($p < 0.001$; Table 2). There were significant differences among regions at the S1 level, with the sacral body, ilium, and sacral ala having 99.1 ± 72.1, 124 ± 90.1, and −10.7 ± 61.9 HU, respectively ($p < 0.001$). There were significant differences among regions at the S2 level, with the sacral body and ilium having HUs of 168 ± 55.4 and 92.8 ± 105, respectively, while the sacral ala
HU Changes After Teriparatide Administration

There was a significant increase in all three lumbar regions following teriparatide administration. The lumbar lamina HU increased 16.4% on average (343 to 400, p < 0.001). The lumbar pedicles HU increased 10.2% on average (219 to 242, p = 0.04), and the lumbar vertebral body HU increased 20.7% (99.1 to 130, p < 0.001; Table 2 and Fig. 3A).

At S1, there was a significant increase in the HUs of the sacral ala (−10.7 to 3.68, p = 0.04), the ilium (124 to 165, p = 0.01), and the sacral body (99.1 to 130, p < 0.05; Table 2 and Fig. 3B). At S2, there was no significant change in the HUs of the sacral ala (−11.9 to −8.43, p = 0.54) or the ilium (92.8 to 112, p = 0.19); however, the HUs of the sacral body significantly increased (168 to 189, p < 0.05). The HUs at the sciatic notch had a significant 86.7% increase after teriparatide treatment (30.6 to 57.1, p = 0.004).

DXA BMD

Based on DXA, femoral neck BMD (g/cm²) increased by 3% (0.65 to 0.66, p = 0.07), lumbar BMD increased by 10% (0.83 to 0.92, p < 0.001), and total hip BMD increased by 5% (0.72 to 0.75, p = 0.001).

HU Correlation to DXA BMD

Prior to teriparatide treatment, the Pearson correlation coefficients between lumbar spine HUs and femoral neck BMD, lumbar BMD, or total hip BMD were 0.27 (25th–75th IQR −0.14 to 0.61, p = 0.20), 0.45 (IQR 0.10 to 0.69, p = 0.01), and 0.46 (IQR −0.03 to 0.77, p = 0.06), respectively (Table 3). The Pearson correlation coefficients between sacropelvic area HUs and the femoral neck BMD, lumbar BMD, or total hip BMD were 0.42 (IQR 0.02 to 0.71, p = 0.04), 0.38 (IQR 0.03 to 0.65, p = 0.04), and 0.49 (IQR 0.02 to 0.79, p < 0.05), respectively.

Following teriparatide treatment, the Pearson correlation coefficients between lumbar spine HUs and femoral neck BMD, lumbar BMD, or total hip BMD were 0.43 (IQR −0.03 to 0.67, p = 0.07), 0.43 (IQR 0.07 to 0.69, p = 0.02), and 0.51 (IQR 0.02 to 0.80, p = 0.04), respectively. The Pearson correlation coefficients between sacropelvic area HUs and the femoral neck BMD, lumbar BMD, or total hip BMD were 0.55 (IQR 0.20 to 0.78, p = 0.004), 0.39 (IQR 0.02 to 0.67, p = 0.04), and 0.66 (IQR 0.25 to 0.87, p = 0.005), respectively. Figure 4 demonstrates representative correlations between lumbar DXA BMD and lumbar HUs and between total hip DXA BMD and sacropelvic HUs, both following treatment with teriparatide, with the highest correlation between total hip DXA and sacropelvic HUs.

Reliability

Overall, the HU measurements had an interrater reliability of 0.88 (p < 0.0001). Lumbar spine measurements had an interrater reliability of 0.99 (p < 0.0001), while

TABLE 1. Overall characteristics of 40 patients receiving teriparatide treatment

| Characteristic                        | Value          |
|---------------------------------------|----------------|
| Average age in yrs (range)            | 67 ± 12 (36–84) |
| Sex: F/M                              | 34:6           |
| Mean height in cm                     | 161 ± 8        |
| Mean weight in kg                     | 62 ± 15        |
| Mean BMI in kg/m²                     | 24 ± 6         |
| Mean duration of teriparatide therapy in mos (range)| 20 ± 6 (6–27) |
| Average time btw pre & post CT scans in mos | 39 ± 21       |
| Average time btw pre & post DXA in mos | 28 ± 13        |
| Baseline mean DXA T-score at femoral neck | −2.6 ± 0.9    |
| Baseline mean DXA T-score at lumbar spine | −2.4 ± 1.5    |
| Baseline mean DXA T-score at total hip | −2.2 ± 1.1     |

Values are expressed as the mean ± SD, unless indicated otherwise.

TABLE 2. Regional HU values pre- and posttreatment with teriparatide

| Region               | Pretreatment HUs | Posttreatment HUs | % Change | p Value |
|----------------------|------------------|------------------|----------|---------|
| Lumbar lamina        | 343              | 400              | 16.4     | <0.001  |
| Lumbar pedicle       | 219              | 242              | 10.2     | 0.04    |
| Lumbar vertebral body| 111              | 134              | 20.7     | <0.001  |
| S1 ilium             | 124              | 165              | 33.1     | 0.01    |
| S1 sacral ala        | −10.7            | 3.68             | NA       | 0.04    |
| S1 vertebral body    | 99.1             | 130              | 23.8     | <0.05   |
| S2 ilium             | 92.8             | 112              | 20.7     | 0.19    |
| S2 sacral ala        | −11.9            | −8.43            | NA       | 0.54    |
| S2 vertebral body    | 168              | 189              | 12.5     | <0.05   |
| Sciatic notch        | 30.6             | 57.1             | 86.7     | 0.004   |

NA = not available, written when either HU value was negative, as this made assessment of percent change difficult to interpret.

Boldface type indicates statistical significance.
sacropelvic measurements had an interrater reliability of 0.78 (p < 0.0001).

**Discussion**

In our study, teriparatide increased average HU values across all regions of the lumbar spine and most regions in the sacropelvic area. Additionally, we noted that compared to the other regions, the vertebral bodies of the lumbar-sacral vertebra showed the most consistent and most robust response to teriparatide. In osteoporosis, there is a loss of overall trabecular bone mass, with a reduced trabecular bone volume fraction and lower trabecular number.29 However, it is unknown why certain bony regions may exhibit a differential response to anabolic agents.30

In our study, the more consistent and seemingly larger increase in response to teriparatide at the vertebral body likely results from the increased ratio of trabecular bone relative to cortical bone in the vertebral body, as teriparatide has a greater effect on trabecular bone than cortical bone.31–33 For instance, both Genant et al.32 and Borggrefe et al.31 reported an increase in femoral trabecular volumetric BMD (vBMD) with a decrease in proximal femoral cortical vBMD at 12 and 24 months, respectively. There is an early transient increase in femoral cortical bone turnover and cortical porosity following initiation of teriparatide therapy, though this may not decrease bone strength.34 Additionally, there is evidence that loading through exercise may augment the response to teriparatide,35 and it has been proposed that an increased catabolic effect of teriparatide on cortical porosity at the tibia and radius may result from different loading conditions.33

Additionally, preclinical and limited clinical data suggest that teriparatide has the potential to increase spinal fusion rates and decrease pseudarthrosis rates.7,21,22,36 For instance, in a rabbit posterolateral fusion model, increasing doses of teriparatide up to 40 μg/kg/day resulted in an increased fusion mass volume as compared to that in the autograft-alone control group.36 In a clinical study, Ohtori et al.7 found that a daily injection of teriparatide following lumbar fusion in osteoporotic women resulted in a significantly lower rate of pedicle screw loosening compared to that in patients taking risedronate and in controls. In a follow-up study, the same group reported an increased fusion rate as well as a faster time to fusion.22 Similarly, Ebata et al.28 found that weekly teriparatide administration following lumbar interbody fusion surgery in patients with osteoporosis increased bone growth within the fusion site at even 4 months postoperatively. Other groups have reported an increased rate of pelvic fracture healing and improved function in elderly osteoporotic women following only 8 weeks of PTH treatment.27 Therefore, it is

| TABLE 3. HU correlation with DXA BMD before and after treatment with teriparatide |
|---------------------------------|------------------|------------------|------------------|
| Level                          | Femoral Neck     | Lumbar           | Total Hip        |
| Lumbar before                  | 0.27 (−0.14 to 0.61, 0.20) | 0.45 (0.10 to 0.69, 0.01) | 0.46 (−0.03 to 0.77, 0.06) |
| Lumbar after                   | 0.37 (−0.03 to 0.67, 0.07) | 0.43 (0.07 to 0.69, 0.02) | 0.51 (0.02 to 0.80, 0.04) |
| Sacropelvic region before      | 0.42 (0.02 to 0.71, 0.04) | 0.38 (0.03 to 0.65, 0.04) | 0.49 (0.02 to 0.79, <0.05) |
| Sacropelvic region after       | 0.55 (0.20 to 0.78, 0.004) | 0.39 (0.02 to 0.67, 0.04) | 0.66 (0.25 to 0.87, 0.005) |

Boldface type indicates statistical significance.
important to better define how teriparatide’s impact can be optimized for a specific surgery involving a given region. A significant degree of regional variation in HUs was observed, especially in the sacrum, where the sacral ala had the worst bone density with negative HU measurements. The negative values suggest red marrow loss with large voids in trabeculae and/or fatty infiltration of the bone. These results were similar to those of a prior study on a general population, in which the central sacral ala had the lowest HU measurements with a mean of 24, while the L1 and S1 vertebral bodies were 165 and 224 HUs, respectively. There was a clear discrepancy in measurements in the sacropelvic area and the lumbar spine overall, with the lumbar spine having significantly higher measurements overall, suggestive of higher BMD. However, in contrast to a prior study assessing sacral HUs, we did not find the S2 level to have less dense bone than the S1 level, which was potentially attributable to differences in populations, with our population being significantly more osteoporotic.

The sacral ala at both S1 and S2 levels had by far the worst baseline HUs, with even negative measurements, and there was no significant improvement in the HUs of the ala and ilium at the S2 level. This result may be attributable to the relatively small sample size and may suggest that teriparatide administration has limited effects at these regions or that these regions had such poor baseline bone density with significant fatty infiltration that even 6 months of teriparatide treatment was not adequate to obtain meaningful improvement. Since both the lumbar spine and pelvis have large stores of cancellous bone, it is still unknown what caused the regional variation detected between the spine and pelvis at baseline. Since reciprocal work has not been performed in subjects with normal bone density, it is not known if this difference is intrinsic or an effect of differential bone loss due to osteoporosis that is greater in the sacrum than in the low lumbar spine. However, these observations are consistent with the fact that the low lumbar spine is less affected by fragility fractures than more cephalad levels, while the sacrum is a common site of insufficiency fractures.

When considering pelvic fixation with S2 alar-iliac (S2AI) screws, our results indicate that even following teriparatide treatment in osteoporotic patients, there may be minimal biomechanical contribution from the sacral fixation with limited benefit on pullout strength, which is integral to durable sacropelvic fixation. Additionally, with a renewed focus on sacroiliac fixation and fusion, these results suggest that fixation in the sacral ala may be limited in the osteoporotic patient population. Alternatively, the findings may emphasize the importance of obtaining bicortical or tricortical fixation of sacral pedicle screws in this population given that the most marked improvements in HUs occurred at the vertebral body. Overall, given the poor bone density in the cancellous bone of the pelvis, trajectories such as S2AI screws that involve multiple cortical surfaces may be superior in osteoporotic patients. However, further biomechanical studies are required to better understand the implications of this regional heterogeneity. To our knowledge, this is one of few studies analyzing changes in HUs throughout the lumbosacral spine and pelvis following teriparatide therapy. Our focus on region-specific changes may better elucidate the mechanistic effects of at least 6 months of perioperative teriparatide administration and eventually permit patient-specific screw trajectory or construct considerations.

Improvement in the BMD was significant on DXA of the femoral neck, lumbar spine, and hip regions, with posttreatment increases of 3%, 10%, and 5%, respectively. However, there was a low to moderate degree of correlation between baseline HUs and DXA BMD, as well as the improvements observed in HUs and DXA BMD. The correlation was stronger after teriparatide administration and tended to be stronger in the sacropelvic region than in the lumbar region, though these were still relatively moderate correlations. Consistent with our recently published findings on the lumbar vertebral body alone, the increase in HUs we detected in this study was, on a percentage basis, out of proportion to the increase observed on DXA scans. This may suggest that HU analysis is more sensitive than DXA scans when assessing teriparatide-related changes to bone density, likely since HU measurements allow for more region-specific measurements than DXA scans, with less interference from degenerative pathology of the spine. There is further support for the use of
CT-based measurements as opposed to DXA, as Graeff et al. found that quantitative CT was better associated with fracture status and severity than DXA measurements at the spine or hip.

There are several limitations to this study, including its retrospective nature and small sample size. Additionally, as with most studies on HU assessments, there can be differences in the measurements based on the use of different CT machines and varied distances from the patient. Furthermore, there can be significant bone loss following the completion of teriparatide treatment, with a recommendation for transitioning to bisphosphonate therapy. Since we did not make all the posttreatment HU measurements while patients were still on teriparatide therapy, it is possible that some of the posttreatment measurements would be artificially low. We were not equipped to determine the optimal duration of teriparatide treatment in this study because of our small sample size and the variable amount of time between cessation of teriparatide treatment and remeasuring with CT scans. However, we generally recommend at least 6 months of teriparatide preoperatively with postoperative continuation to complete the full 24 months of treatment. Finally, we realize that BMD assessed by any technique is merely a proxy for bone strength and bony fusion potential; therefore, further studies correlating higher HU values and improved clinical outcomes are needed.

Conclusions
There was significant regional variation in lumbar and sacropelvic HUs, with most regions significantly increasing after at least 6 months of teriparatide treatment. The sacropelvic area had lower HU values—with even negative values of HUs in the sacral ala—more regional variation, and a higher degree of correlation with BMD as measured on DXA. While teriparatide treatment resulted in HUs > 110 in most of the lumbosacral spine, the sacral ala remained with negative HUs suggestive of severe osteoporosis, which may limit the effectiveness of fixation in this region.

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Disclosures
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Conception and design: Elder, Flanigan, Mikula, Peters, Oushy. Acquisition of data: Elder, Flanigan, Mikula, Peters, Oushy. Analysis and interpretation of data: Elder, Flanigan, Mikula, Peters, Oushy, Anderson. Drafting the article: Elder, Flanigan, Mikula, Fogelson, Bydon, Freedman, Sebastian, Currier, Nassr, Kennel, Anderson, Polly. Critically revising the article: Elder, Flanigan, Mikula, Oushy, Fogelson, Bydon, Freedman, Sebastian, Currier, Nassr, Kennel, Polly. Approved the final version of the manuscript on behalf of all authors: Elder. Statistical analysis: Flanigan, Mikula, Oushy. Administrative/technical/material support: Elder. Study supervision: Elder, Flanigan, Mikula, Fogelson, Bydon, Freedman, Sebastian, Currier, Nassr, Kennel, Polly.

Supplemental Information
Previous Presentations
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