Cortical Porosity Identifies Women with Osteopenia at Increased Risk for Forearm Fractures†

Short Title: Cortical Porosity and Bone Fragility

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Abstract

Background Most fragility fractures arise among the many women with osteopenia, not the smaller number with osteoporosis at high risk for fracture. Thus, most women at risk for fracture assessed only by measuring areal bone mineral density (aBMD) will remain untreated.

Methods We measured cortical porosity and trabecular bone volume/total volume (BV/TV) of the ultradistal radius (UDR) using high-resolution peripheral quantitative computed tomography, aBMD using densitometry, and 10-year fracture probability using the country-specific FRAX tool in 68 postmenopausal women with forearm fractures and 70 age-matched community controls in Olmsted County, Minnesota.

Results Women with forearm fractures had 0.4 standard deviations (SD) higher cortical porosity and 0.6 SD lower trabecular BV/TV. Compact-appearing cortical porosity predicted fracture independent of aBMD; odds ratio [OR] 1.92 (95%CI, 1.10-3.33). In women with osteoporosis at the UDR, cortical porosity did not distinguish those with, from those without, fractures because high porosity was present in 92% and 86% of each group respectively. By contrast, in women with osteopenia at the UDR, high porosity of the compact-appearing cortex conferred an OR for fracture of 4.00 (95%CI, 1.15-13.90).

Conclusion In women with osteoporosis, porosity is captured by aBMD and so measuring UDR cortical porosity does not improve diagnostic sensitivity. However, in women with osteopenia, cortical porosity was associated with forearm fractures.

Key Words bone mineral density, cortical porosity, forearm fractures, microarchitecture, trabecular bone.
Introduction

The terms ‘osteoporosis’ and ‘osteopenia’ were originally coined to convey the notion that an individual is susceptible to sustaining a fracture following minimal trauma because there is ‘not enough bone’\(^1-3\). To formalize this imprecise notation, a working group of the World Health Organization (WHO) defined osteoporosis as a “systemic skeletal disease characterized by low bone mass and micro-architectural deterioration of bone tissue with a consequent increase in bone fragility and susceptibility to fracture”\(^4\). ‘Osteoporosis’ was quantified as a femoral neck (FN) aBMD T-score of -2.5 SD or more below the young normal mean, osteopenia as a T-score between -1 and -2.5 SD, and normal aBMD as a T-score above -1 SD\(^5\).

Subsequent epidemiological research confirmed that fracture risk increases as aBMD decreases. However, these studies also showed that ~50% of all fractures, ~50% all recurrent fractures, and much of the accompanying morbidity, mortality and costs to the community arise from the large proportion of the population with osteopenia at modest risk for fracture, not the smaller fraction with osteoporosis at high risk for fracture\(^6-9\). This observation identified a challenge to curbing the population burden of fractures.

Among the large group of women with osteopenia, there exists a substantial subgroup with bone fragility contributing to the burden of fractures. If an aBMD measurement alone is used in an osteoporosis screening program, women with osteopenia will be excluded from further investigation and so will not be offered treatment\(^10-12\). One approach to case finding - identifying those at risk for fracture in need of treatment is the use of the fracture risk assessment tool (FRAX)\(^11,12\). Another approach is to identify the structural basis of the bone fragility not captured by the aBMD measurement and thereby to quantify “microarchitectural deterioration of bone tissue”, the descriptive component of the definition of ‘osteoporosis’.

Trabecular bone loss and vertebral fractures are historical hallmarks of osteoporosis. However, ~80% of the skeleton is cortical; 80% of all fractures are non-vertebral and 30% of these are forearm fractures\(^13\). Moreover, about 70% of all the appendicular bone lost during aging is cortical and results from intracortical remodelling which occurs throughout the cortex but is particularly vigorous in the cortico-trabecular junctional (transitional) zone where the cortical and trabecular compartments merge (figure 1)\(^14\). Remodelling during advancing age becomes unbalanced and removes more bone than it deposits leaving residual cortical porosity, which increases bone fragility exponentially and is a quantifiable ‘footprint’ of bone loss\(^14-16\).

Quantifying porosity \textit{in vivo} has become possible with the recent development of high-resolution peripheral quantitative computed tomography (HRpQCT), a noninvasive method of image acquisition, and StrAx1.0, a new method of image analysis that permits quantification of porosity, even porosity due to pores under 100 microns (the diameter of >80% of cortical pores), and quantifies porosity of the transitional zone\(^14,17\). The aim
of this study was to determine (i) whether bone microarchitecture, particularly cortical porosity, predicts fracture; (ii) whether porosity does so independent of aBMD and FRAX; and (iii) whether combining a measurement of forearm microarchitecture and aBMD (at the forearm or femoral neck) identifies more women with fractures than aBMD alone.

**Methods**

**Participants**

As previously reported, 100 postmenopausal women aged ≥50 years with a distal forearm fracture were matched with 105 controls from an age-stratified random sample of women from Olmsted County. The fracture occurred 7 (3-13) months (median (IQR)) before the investigation. Fragility fracture was defined on the basis of the original description of the fall that led to fracture using the classification of Palvanen et al. This corresponds to the convention of a moderate trauma distal forearm fracture resulting from a fall from standing height or less. Controls had no history of a fracture after 35 years of age. Of the subjects analyzed here, 21/68 (31%) of cases and 21/70 (30%) of controls received bisphosphonate or estrogen therapy. Observations were reported in the whole group previously, and here were no different after excluding treated subjects. The cohort was >96% Caucasian. The study was approved by the Mayo Clinic Institutional Review Board and the present analysis was based on de-identified data.

**Measurement of micro-architecture, aBMD and FRAX score**

Microarchitecture was assessed at the non-fractured ultradistal radius (UDR) using HRpQCT (XtremeCT, Scanco Medical AG, Brüttisellen, Switzerland). Quality control was monitored by daily scans of phantoms (hydroxyapatite (HA) rods, QRM, Moehrendorf, Germany). Movement artefacts precluded use of scans in 32 cases and 35 controls leaving 68 cases (mean age ± SD, 63 ± 9 years) and 70 controls (mean age 66 ± 10 years; p = 0.09). Age and aBMD did not differ between excluded and included subjects. The coefficient of variation calculated for the same patients at different times after repositioning was < 1.5% for volumetric density and 0.54 to 3.98% for cortical porosity.

StrAx1.0 is a new algorithm that segments cortical bone from background and from the trabecular compartment, into its compact appearing, outer and inner transitional zones and in so doing correctly assigns the trabecularized cortex (i.e., cortical fragments) to the transitional zone rather than to the medullary compartment which results in overestimation of ‘trabecular’ density. The outer transitional zone is defined as the trabecularized cortex that is adjacent to the compact-appearing cortex whereas the inner transitional zone is the trabecularized cortex adjacent to the medullary cavity. The latter also contains some true trabecular bone. The 40 most proximal slices are analyzed using ~3600 radial attenuation profile curves around each slice. All voxels within the periosteal envelope are analyzed and most are composite voxels as they contain both void volume and mineralized bone matrix volume. The proportions of each are quantified using an interpolation
function derived from voxels with zero attenuation (porosity). Voxels with attenuation produced by fully mineralized bone (density of 1200 mgHA/cc or greater) are assigned 100%. The void volume of a voxel = 100 - mineralized bone volume fraction. Total porosity is the average of the summed void volume fractions of all voxels producing an attenuation ≤ 80% than produced by fully mineralized bone (≥1200 mgHA/cc). Attenuation between 80-100% of this maximum can be produced by either heterogeneity in bone tissue mineralization density or a small pore (≤ ~35 µm diameter) in a voxel containing fully mineralized bone matrix. Thus, porosity may be underestimated by erroneously discarding existing pores of ≤ 35µm diameter. UDR and femoral neck (FN) aBMD (g/cm²) were measured by DXA (Lunar Prodigy, GE Healthcare, USA).

The 10-year probability of a major fracture (hip, clinical spine, proximal humerus or forearm fracture) and of a hip fracture were assessed using the country-specific WHO risk assessment tool FRAX (version 3.8), including FN aBMD 22. Information on the FRAX input variables were available in all women. The forearm fracture was not included as the aim was to estimate the 10-year probability of the fracture before the event, not the probability of fracture after this event.

Statistical analysis

Fracture risk associated with aBMD and microstructure was estimated by odds ratios (OR) from logistic regression models adjusting for age, FRAX probabilities, UDR and FN aBMD. Group differences were summarized using the ΔSD, the mean value in cases standardized using the mean and SD in controls. Linear regression was used to study the relationship between microstructure and age and to test for differences in slopes and intercepts between cases and controls. The areas under a receiver operating curves (AUC) were compared using nonparametric methods 23,24. Cases and controls were pooled and partitioned into those with osteoporosis, osteopenia or normal aBMD at the UDR and FN. ORs for fracture were computed defining ‘high’ cortical porosity (>90th centile) and ‘low’ trabecular BV/TV (<10th centile) in 40 healthy premenopausal women in Melbourne (age 27 years, range 21-31). A p < 0.05 (two tailed) denoted statistical significance.

Results

Cortical porosity predicts forearm fractures independently of forearm aBMD

Women with forearm fractures had higher porosity of the compact-appearing cortex, outer and inner transitional zones, and lower trabecular BV/TV than controls (Table 1). Porosity increased linearly as age advanced with no difference in the slopes of the regression lines but the y-intercepts (porosity) were 0.35, 0.46 and 0.61 SD higher in the respective cortical compartments in women with fractures than controls (all p <0.001) (Figure 2).

Porosity of each cortical compartment and trabecular BV/TV predicted forearm fractures. Porosity of the compact appearing cortex and outer transitional zone, but not inner transitional zone or trabecular BV/TV,
predicted fracture independently of UDR aBMD. Porosity discriminated cases from controls with AUCs ranging from 0.71 to 0.83 (all AUCs > 0.5 with p < 0.001) (Table 1).

**Cortical porosity is captured by the diagnostic threshold of -2.5 SD for osteoporosis**

Osteoporosis of the UDR was present in almost twice the number of cases than controls; 26 (38%) cases and 14 (20%) controls conferring an OR of 2.48 (95% CI, 1.15-5.31). High porosity in one or more cortical compartments was common in women with osteoporosis; 92% with a fracture and 86% without a fracture. Thus, porosity was captured by the diagnostic category of ‘osteoporosis’ at the UDR. Adding of a measure of porosity to aBMD did not identify more women with osteoporosis with fractures than aBMD alone.

**Cortical porosity increases the diagnostic sensitivity of osteopenia, not normal aBMD**

Osteopenia of the UDR was present in 26 (38%) cases and controls 25 (36%). Osteopenia alone was not associated with fracture as the OR was 1.11 (95% CI, 0.56-2.22) (Figure 3). However, over twice the number of women with osteopenia and a fracture had high porosity of the compact appearing cortex than did women with osteopenia without a fracture (50% versus 20% respectively), conferring an OR of 4.00 (95% CI, 1.15-13.90) and a specificity of 80%. Porosity of the outer and inner transitional zones conferred ORs of 3.17 (95% CI, 0.96-10.48) and 2.93 (95% CI, 0.84-10.25), respectively.

Normal UDR aBMD was present in 16 (24%) cases and 31 (44%) controls and was protective, with an OR of 0.39 (95% CI, 0.19-0.81). High porosity was not more prevalent in those with, than without a fracture. Thus, adding a measure of porosity did not identify more women with normal aBMD with a fracture.

**Cortical porosity, WHO diagnostic categories using FN aBMD, and forearm fracture risk**

Compact-appearing and outer transitional zone porosity of the UDR predicted forearm fractures independently of FN aBMD (Table 1). Only 9% of women with forearm fractures had FN osteoporosis; 72% had FN osteopenia and 19% had normal FN aBMD. No significant associations with fracture were detected with FN osteoporosis alone [OR 2.16 (95% CI, 0.52-9.01)] or FN osteopenia alone [1.62 (95% CI, 0.79-3.31)]. Moreover, FN osteopenia plus high porosity of the UDR inner transitional zone, or FN osteopenia plus low UDR trabecular BV/TV were each associated with forearm fracture, conferring ORs of 2.92 (95% CI, 1.18-7.20) and 2.44 (95% CI, 1.01-5.92) respectively. Normal FN aBMD was protective with an OR 0.45 (95% CI, 0.21-0.99).

**Cortical porosity predicts forearm fractures independently of FRAX**

The ten-year probability of a major osteoporotic or hip fracture did not differ significantly between cases and controls (Table 1). By contrast, there were significant correlations between the probability of a major fracture and cortical porosity (r ranging from 0.26 to 0.40, all p<0.001). Compact-appearing cortex, outer and inner
transitional zone porosities remained significantly associated with forearm fracture after adjustment for FRAX, with ORs of 2.11 (95% CI, 1.35-3.30), 2.06 (95% CI, 1.32-3.22) and 2.48 (95% CI, 1.65-3.73), respectively.

Discussion

We report the following: (i) women with forearm fractures had micro-architectural deterioration; cortical porosity was increased and trabecular bone volume fraction was reduced. (ii) Both predicted forearm fractures, but only cortical porosity did so independently of UDR or FN aBMD. (iii) The diagnostic threshold for ‘osteoporosis’ (T-score ≤ -2.5 SD), whether based on UDR or FN aBMD, captured high cortical porosity and low trabecular bone volume of the UDR whether a forearm fracture was present or not. Thus, after finding a BMD T-score ≤ -2.5 at the UDR or FN, measuring microstructure at the UDR did not identify more women with forearm fractures than measuring aBMD alone. These observations support the inclusion of “microarchitectural deterioration” in the WHO definition of osteoporosis.

While the diagnostic category of ‘osteoporosis’ captures microarchitectural deterioration in women at high risk for fracture, most fractures in the community arise from the larger segment of the population with an aBMD T-score less severely reduced than -2.5 SD. In this study, at the UDR, 38% of the women with a forearm fracture had osteopenia, while 24% had normal aBMD. At the FN, 72% had osteopenia and 19% had normal aBMD. As a group, these women were not at increased risk for fracture since neither UDR nor FN osteopenia alone was associated with fracture. Thus, physicians finding a UDR or FN aBMD T-score in the osteopenic range will be disinclined to initiate treatment even though most forearm fractures arise from this group.

Combining risk factors, as used in FRAX, assists in identifying individuals in need of treatment and avoids treating individuals at low risk for morbid events. Similarly, by adding a measurement of cortical porosity at the UDR, we identified a subset of women with UDR or FN osteopenia contributing to the burden of fractures. Thus, in women found to have osteopenia (the usual outcome in the community), the data suggest that it may be appropriate to also measure UDR porosity. Finding high porosity, which compromises bone strength out of proportion to the modest reduction in aBMD that characterizes osteopenia, identifies individuals in need of treatment who would not be identified otherwise.

Women with normal UDR or FN aBMD were relatively protected against fracture. Nevertheless, 24% of all women with forearm fractures had normal UDR aBMD. Measurement of porosity did not distinguish women with normal UDR aBMD with a fracture from those without a fracture because the prevalence of microarchitectural abnormalities was similar in those with and without fractures. In these women, the fracture may have been the result of more severe trauma than was estimated using the semiquantitative classification of trauma severity, or may be due to the result of abnormalities in bone material composition and structure yet to be identified.
Porosity throughout the cortex was associated with fracture, but only porosity of the compact-appearing cortex and outer transitional zone, not porosity of the inner transitional zone, predicted fracture independently of aBMD. This site-specific independence from aBMD may be due to the location of the porosity. While porosity of the inner transitional zone is larger than that of the compact appearing cortex and outer transitional zone, it is positioned nearer the neutral axis adjacent to the medullary canal. Deficits in mineralized bone matrix volume at this location may contribute less to loss of bending strength than deficits in mineralized bone matrix volume positioned further from the neutral axis.

Cortical porosity in adulthood is the net result of ‘peak’ porosity achieved during growth, constituted mainly by the Haversian and Volkmann canals, and the subsequent increase in porosity produced by age-related intracortical remodeling upon these canals which enlarges them focally and produces coalescent and giant pores in cross section as age advances. In this study, porosity increased across age, but the slope of porosity as a function of age was no greater in women with, than without, fractures, suggesting that women sustaining a fracture may assemble a bone with a higher peak porosity.

Porosity measured in this study was several-fold higher than the 1-16% porosity reported using HR-pQCT. While cortical bone is ‘compact’ relative to trabecular bone, the term ‘compact’ is a misnomer. Cortical bone is a three dimensional structure comprising mineralized bone matrix volume and a void volume formed at the nano-scale level by voids within and between collagen fibrils, and formed at the micro-scale level by the osteocyte lacunar-canalicular system and by the Haversian and Volkmann canals.

These voids are not ‘empty’; they are fluid filled and so the measurement of bone water content provides an accurate measurement of the component of cortical bone that is void volume. Direct measurements of cortical bone water content across species using deuterium oxide or dehydration experiments report a void volume ranging from 15 to 40%. Synchrotron radiation based µCT assessment of histomorphometric specimens suggests that the diameter of ~60% of pores is < 90 µm; ~20% is 90-180 µm and ~20% is > 180 µm. The values for porosity reported here are in agreement with the above and are compatible with the physiological role of the canals which house bone’s vasculature.

We suggest that the low porosities reported in previous studies are incompatible with the direct measurements cited above and are incompatible with the provision of an effective vascular supply since bone is a highly vascularized organ and receives 10 to 20% of total cardiac output essential for nutrient transport and waste removal. The discrepancies are likely to be the result of differences in image analysis techniques. While HR-pQCT has a voxel size of 82 µm, its in-plane resolution is about 130 µm. This precludes the quantification of pores lower than this value. As reported above, more than 60% of pores are under 100 microns. Excluding these, results in underestimation of porosity which depends upon the cortical compartment (compact-appearing
or transitional zone) measured, age, sex, and the underlying disease present $^{17,38}$. Indeed, the larger pores may contribute disproportionately more to total porosity than smaller pores even though smaller pores are more numerous in samples from elderly in whom cortex is porous and fragmented. However, in samples from younger individuals, with a cortex mostly compact in appearance and not fragmented, the porosity created by smaller pores is likely to account for a larger proportion of the total porosity. The magnitude of the underestimation in each of these states will be the subject of future research.

This study has several limitations. The study was cross-sectional so determining the relative contributions of peak porosity and age-related bone loss to fracture risk was not possible. Moreover, the sample size may have been too small to detect an increased risk of fracture in some subgroups. About 33% of the patients enrolled in the study were excluded because of motion artifacts, a recognized problem being addressed by shorter scanning times and better forearm immobilization methods in newer imaging devices $^{20}$. Moreover, while the technique used to estimate porosity account for pores smaller than 100 µm, we acknowledge that it misses pore smaller than 35 µm.

In summary and conclusion, postmenopausal women with distal forearm fractures have microstructural deterioration characterized by high cortical porosity and reduced trabecular BV/TV. UDR aBMD in the osteoporosis range (T-score $\leq -2.5D$) captured these abnormalities so measurement of microarchitecture did not identify a greater proportion of women with fracture than did areal BMD in this range. In women with osteopenia, the source of over 50% of all fractures, fracture risk was increased if high porosity was present so measuring porosity improved identification of women with osteopenia with forearm fractures. Thus, measuring porosity is likely to be clinically useful in identifying women at risk for fracture considered at low risk based on their aBMD measurement alone. Further research will be needed to determine whether assessment of microstructure at the spine and proximal femur improves the sensitivity and specificity in identification of women at risk for fracture at those locations.

Disclosures:
AGZ is one of the inventor of StrAx 1.0. ES & RZ are inventors of StrAx 1.0 algorithm and directors of Straxcorp. StrAx software is now being actively marketed. RZ has received Consulting fees / lecture fees / grant support from Amgen, MSD, Servier. JK has received Consulting fees / lecture fees / grant support from Amgen, D3A, GSK, Lilly, Medimaps, Merck. ES have received Consulting fees / lecture fees / grant support from Amgen, MSD, Novartis, Sanofi Aventis, Servier. RZ and ES are inventors of the Strax method and directors of StrAxcorp. Other Authors have no disclosures to declare.
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Author’s roles: Drafting manuscript: YB, ES, SK, JK, JM. Revising manuscript content: YB, ES, EA, RZ JP, JM, JK and SK. Approving final version of manuscript: YB, RZ, EA, AGZ, JP, AB, JM, HJ, JK, SK and ES. ES takes responsibility for the integrity of the data analysis.
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Tables & figures legend:

Table 1  
Comparison of areal bone mineral density (aBMD) at the ultradistal radius (UDR) and femoral neck (FN), FRAX score, cortical porosity and trabecular bone volume fraction (BV/TV) between postmenopausal women with (cases) or without forearm fracture (controls). Odds ratio (OR) and 95% confidence intervals (95% CI) are reported for 1 standard deviation (SD) increment. Area under the receiving operating curve (AUC) represents the ability of the variables to discriminate the fracture status.

Figure 1  
Representative segmented image obtained at the ultradistal radius using non threshold-based image analysis in a postmenopausal women with (Case) and without (Control) forearm fracture. The full cross section and the magnified image show the presence of porosity within the compact appearing cortex (green) and the outer (white) and inner (red) transitional zones, and loss of trabecular bone (yellow) in the case, and less so in the control.

Figure 2  
Cortical porosity in the compact-appearing cortex, outer transitional zone, inner transitional zone and trabecular bone volume fraction (BV/TV) versus age in women with distal forearm fractures (Cases, black dots, solid line) and without forearm fracture (controls, white open circles, dotted line).

Figure 3  
Odds ratio (OR) and 95% confidence interval (95% CI) for distal fracture associated with areal bone mineral density (aBMD) alone and in combination with high cortical porosity assessed in the compact appearing cortex in women with osteopenia at the ultradistal radius. \(^a\)p < 0.05.
<table>
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<tr>
<th>Variable</th>
<th>Cases (n=68)</th>
<th>Controls (n=70)</th>
<th>Δ SD (95% CI)</th>
<th>unadjusted OR for fracture (95% CI)</th>
<th>adjusted for UDR aBMD OR for fracture (95% CI)</th>
<th>adjusted for FN aBMD OR for fracture (95% CI)</th>
<th>AUC (95% CI)</th>
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<td><strong>Ankle BMD</strong></td>
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<td>UDR (g/cm²)</td>
<td>0.37 (0.08)</td>
<td>0.42 (0.07)</td>
<td>-0.51 (-0.62; -0.21)</td>
<td>2.46 (1.57; 3.80) **</td>
<td>-</td>
<td>-</td>
<td>0.72 (0.63; 0.80)</td>
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<td>FN (g/cm²)</td>
<td>0.81 (0.10)</td>
<td>0.86 (0.14)</td>
<td>-0.53 (-0.64; -0.23)</td>
<td>2.32 (1.48; 3.55) **</td>
<td>-</td>
<td>-</td>
<td>0.71 (0.62; 0.79)</td>
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<td><strong>FRAX score</strong></td>
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<td>Major fracture (%)</td>
<td>12.51 (7.82)</td>
<td>11.95 (6.03)</td>
<td>-0.07 (-0.17; 0.10)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.53 (0.44; 0.63)</td>
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<td>Hip fracture (%)</td>
<td>2.84 (4.93)</td>
<td>2.70 (5.31)</td>
<td>0.02 (-0.20; 0.25)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.57 (0.47; 0.67)</td>
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<td><strong>Poroetry</strong></td>
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<td>Compact-appearing cortex (%)</td>
<td>36.07 (7.86)</td>
<td>26.02 (7.62)</td>
<td>0.40 (0.07; 0.73)</td>
<td>2.59 (1.56; 4.33) **</td>
<td>1.92 (1.10; 3.33)  *</td>
<td>2.25 (1.33; 3.82)  *</td>
<td>0.72 (0.64; 0.81)</td>
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<tr>
<td>Outer transitional zone (%)</td>
<td>39.19 (5.53)</td>
<td>39.06 (5.95)</td>
<td>0.03 (0.05; 0.71)</td>
<td>2.39 (1.47; 3.99) **</td>
<td>1.75 (1.02; 2.96)  *</td>
<td>2.01 (1.20; 3.34)  *</td>
<td>0.72 (0.63; 0.80)</td>
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<td>Poroetry inner transitional zone (%)</td>
<td>81.37 (3.15)</td>
<td>79.36 (3.41)</td>
<td>0.51 (0.24; 0.84)</td>
<td>2.11 (1.41; 3.14) **</td>
<td>1.38 (0.81; 2.38)  *</td>
<td>1.75 (1.14; 2.66)  *</td>
<td>0.71 (0.63; 0.80)</td>
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<td>Trabecular BV/TV (%)</td>
<td>1.80 (1.48)</td>
<td>2.64 (1.62)</td>
<td>-0.58 (-0.89; -0.24)</td>
<td>1.79 (1.21; 2.63) **</td>
<td>1.04 (0.62; 1.74)  *</td>
<td>1.43 (0.95; 2.17)  *</td>
<td>0.67 (0.58; 0.76)</td>
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* P<0.05, ** P<0.01, *** P<0.001

P<0.05 vs. controls adjusted for aBMD at UDR
Figure 1
Figure 2
Figure 3

<table>
<thead>
<tr>
<th>Condition</th>
<th>OR (95% CI)</th>
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<tr>
<td>aBMD alone</td>
<td>0.39 (0.19-0.81)</td>
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<td>aBMD + high porosity</td>
<td>1.73 (0.39-7.63)</td>
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<tr>
<td>Osteopenia</td>
<td>1.11 (0.56-2.22)</td>
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<td>4.00 (1.15-13.90)</td>
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</table>
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