The first sacral vertebra (S1) is always chosen as the caudal level for spinal fusion with instrumentation for lumbar degenerative diseases because the L5–S1 level has the highest prevalence of disc degeneration in the lumbar spine, which may require decompression and fusion. In addition, long fusion to S1 for lumbar deformity is associated with better maintenance of sagittal alignment and a lower rate of subsequent disc degeneration of L5–S1. However, fusion to S1 can also have a high rate of complications such as pedicle screw loosening or pseudarthrosis, especially for older patients with osteoporosis.

An S1 body with poor bone stock can cause screw loosening in the bone-implant interface, leading to the failure of fixation. Therefore, preoperative measurement of the bone mineral density (BMD) of the S1 body is essential for surgical planning, which helps surgeons decide whether or not to use other techniques to prevent failure at S1. Dual-energy x-ray absorptiometry (DXA) is a widely used method to evaluate BMD and diagnose osteoporosis, but it is routinely used to measure BMD of the first to fourth lumbar vertebrae (L1–4) and hips. Based on the positive correlation between the BMD of different sites in the same patient, the BMD of the S1 body can usually only be estimated via DXA rather than measured directly. Moreover, DXA can overestimate the BMD of the lumbar spine and cause a missed diagnosis of osteoporosis in lumbar degenerative disease, especially for degenerative lumbar scoliosis, which is a more common disease needing fusion to S1.

Since 2011, the method of using Hounsfield units (HUs)
from clinical CT scanning has been widely recommended to assess BMD and screen for osteoporosis.\textsuperscript{1,3,13,14,17} HU values of lumbar vertebrae from L1 to L5 have excellent reliability, significant correlation with DXA T-scores, and good performance in diagnosing osteoporosis. Observers are able to reduce the effect of lumbar degenerative changes on HU values by manually choosing appropriate regions of interest (ROIs) for measurement. Since lumbar CT is a common examination performed for patients requiring surgery for lumbar degenerative diseases, HUs can be used to evaluate BMD at no extra cost and radiation. There have been a few studies involving the manual measurement of HUs of the S1 body from clinical CT scans,\textsuperscript{5,6,16} which support the use of HUs to determine sacral BMD. However, the correlation of HUs of the S1 body with DXA T-scores and the diagnosis of osteoporosis has not been explored. The present study aimed to evaluate the use of HUs of the S1 body to diagnose osteoporosis and to establish HU thresholds if possible.

**Methods**

**Patients**

We reviewed the records of consecutive patients requiring lumbar surgery for lumbar degenerative diseases at our spine center for the period between July 1, 2015, and December 31, 2015. Inclusion criteria were 1) an age of 50 years or older and 2) lumbar CT scanning and central DXA scanning within a month before surgery. Exclusion criteria were 1) a history of spinal surgery; 2) the presence of spinal tuberculosis, tumor, ankylosing spondylitis, diffuse idiopathic skeletal hyperostosis, and other diseases that could significantly affect bone metabolism and BMD; and 3) the presence of severe degenerative changes in the intervertebral disc or endplate at L5–S1 with no appropriate place for measuring HUs of the trabecular bone in the S1 body.

**BMD Evaluation**

Preoperative lumbar 3D reconstructive CT (tube voltage 120 kV, DEFINITION, Siemens) and picture archiving and communication system (PACS) were used to calculate HU values. Patients with transitional vertebrae were not excluded. The L1 vertebral level was identified as the first non–rib-bearing vertebra, and the level of S1 was counted from L1. HU values of the L1 body were measured according to the method of Pickhardt et al.\textsuperscript{13} As for S1, a circular ROI was placed over the midaxial and midsagittal images of the S1 body, respectively (Fig. 1). The diameter of the axial ROI was half of the anteroposterior diameter of the S1 body. The sagittal ROI was placed in the center of the midsagittal image in B. The rule in placing the sagittal ROI was including as much trabecular bone as possible and avoiding cortical bone and heterogeneous areas.

Statistical analysis was conducted using SPSS software (version 20, IBM Corp.). The intraclass correlation coefficient (ICC) was used to evaluate interobserver and intraobserver reliability (ICC ≥ 0.8 was considered to indicate excellent reliability). The correlation between HU values and T-scores was calculated with the Pearson correlation coefficient. The paired t-test was used for comparing the HU values of different vertebral bodies. Receiver operating characteristic curve (ROC) analysis and area under the curve (AUC) were used to evaluate the values of HU in diagnosing osteoporosis. HU thresholds for diagnosing osteoporosis were adjusted to the nearest integer for the ease of clinical use.

**Results**

A total of 316 patients (143 males and 173 females) were included in this study, and the mean patient age was 61.5 ±
6.9 years old (range 50–81 years). Interobserver reliability in measuring axial HU and sagittal HU was excellent with ICCs of 0.971 and 0.967, respectively. Intraobserver reliability in measuring axial HU and sagittal HU was also excellent with ICCs of 0.990 and 0.983, respectively. The correlations among axial HU, sagittal HU, L1 HU, and average T-score of L1–4 are shown in Table 1. The sagittal HU was lower than axial HU (185.8 ± 59.2 HU vs 206.0 ± 60.5 HU, p < 0.001). The L1 HU was lower than sagittal HU (124.2 ± 38.5 HU vs 185.8 ± 59.2 HU, p < 0.001).

The prevalence of osteoporosis identified by DXA and L1 HU ≤ 110 HU was 50.9% (161/316). The AUCs for using axial HU and sagittal HU to distinguish these osteoporotic patients from non-osteoporotic patients were 0.86 (95% CI 0.82–0.90) and 0.88 (95% CI 0.84–0.92), respectively (Fig. 2). The axial HU threshold of 202 HU had balanced sensitivity and specificity to diagnose osteoporosis (sensitivity: 76%; specificity: 76%). The axial HU thresholds with high sensitivity (90%) or high specificity (90%) were 222 HU and 183 HU, respectively. The sagittal HU threshold of 185 HU had balanced sensitivity and specificity to diagnose osteoporosis (sensitivity: 80%; specificity: 80%). The sagittal HU thresholds with high sensitivity (90%) or high specificity (90%) were 200 HU and 160 HU, respectively.

When we only used DXA to identify osteoporotic patients, the prevalence of osteoporosis was 40.5% (128/316), and the AUCs for using axial HU and sagittal HU to distinguish these osteoporotic patients were also over 0.8 (axial HU: 0.82, 95% CI 0.77–0.87; sagittal HU: 0.83, 95% CI 0.78–0.87).

**Discussion**

Results of this study showed that both axial HU and sagittal HU of the S1 body had positive correlations with DXA T-scores and good performance in diagnosing osteoporosis in patients with lumbar degenerative diseases. The axial HU threshold of 202 HU and sagittal HU threshold of 185 HU had balanced sensitivity and specificity over 75% for distinguishing osteoporosis from non-osteoporosis. Two examples are given to show how to use these HU thresholds to detect osteoporosis (Fig. 3).

Similar to the methods used in previous studies, the ROIs in our study were manually placed in the center of midaxial and midsagittal images of the S1 bodies. The reliability of our measurements was excellent with ICCs comparable to those in the study by Katsuura et al. To our knowledge, this is the first study to show the significant correlation between HUs of the S1 body and DXA T-scores, and we found that their Pearson correlation coefficients were comparable to those between HUs of L1–5 and DXA T-scores in other studies. However, these coefficients of 0.6–0.7 were relatively small, which may be the result of choosing patients with older ages and lumbar degenerative diseases. According to Choi et al., the Pearson correlation coefficients could increase to 0.7 if they had only analyzed the data of patients with mild lumbar degeneration.

There was a significant difference among HUs of different sites: L1 HU < sagittal HU of S1 body < axial HU of S1 body. Hoel et al. also found that HU of L1 (165 HU) was lower than HU of S1 body (220 HU). This implies that we need different thresholds for osteoporosis to assess osteoporosis identified by DXA or L1 HU ≤ 110 HU from non-osteoporosis with axial HU and sagittal HU, respectively.

![FIG. 2. Receiver operating characteristic curves for distinguishing osteoporosis identified by DXA or L1 HU ≤ 110 HU from non-osteoporosis with axial HU and sagittal HU, respectively.](image)

![FIG. 3. Clinical examples of using HU thresholds of the S1 body to detect osteoporosis. Images (A and B) from a 55-year-old female with normal BMD; T-scores of 0.5 at L1, −0.3 at L2, 0.5 at L3, −0.2 at L4, −1 at the femoral neck, and 0.2 at the total hip; and an L1 at 197 HU. Her axial HU (A) and sagittal HU (B) were 311.7 and 284.7, respectively. According to our thresholds with balanced sensitivity and specificity, she was diagnosed with non-osteoporosis. Images (C and D) from a 67-year-old female with osteoporosis; T-scores of −2.1 at L1, −3.2 at L2, −1.1 at L3, −1.6 at L4, −2.7 at the femoral neck, and −2.1 at total hip; and an L1 at 78.3 HU. Her axial HU (C) and sagittal HU (D) were 78.3 and 121.2, respectively. According to our thresholds with balanced sensitivity and specificity, she was diagnosed with osteoporosis.](image)
the BMD of different sites. For example, an average HU value of 175 HU for L1–5 is highly suggestive of normal BMD, but an HU of 175 HU for the S1 body may suggest osteoporosis.

Because of the effect of lumbar degenerative changes on DXA measurements, the diagnosis determined by DXA may not be accurate. Thus, we used the threshold of L1 HU to reduce missed diagnoses of osteoporosis as much as possible. The level of L1 has the lowest rate of degeneration when compared with lower levels (L2–S1) and was recommended for measuring HUs because it can be easily identified on different kinds of CT scans. The results showed that the prevalence of osteoporosis and the AUCs were higher when we combined the criterion of DXA and L1 HU ≤ 110 HU instead of using the criterion of DXA alone. In both ways, we found that the use of axial HU and sagittal HU of the S1 body to diagnose osteoporosis was excellent with AUCs over 0.8, which were also comparable to the AUCs (0.8–0.9) of using HU of L1–L5 to diagnose osteoporosis. To provide more information about HU of the S1 body for clinical use, we identified the HU thresholds of different sensitivities and specificities. An HU threshold of balanced sensitivity and specificity may give general information about the correlation between HU and the diagnosis of osteoporosis. An HU threshold of high sensitivity can identify patients who need further evaluation of BMD. Furthermore, patients meeting the HU threshold of high specificity are very likely to have an osteoporotic S1 body despite the coexistence of relatively normal T-scores or HU values for L1–5. In line with the order of HU values of different sites (L1 HU < sagittal HU of S1 body < axial HU of S1 body), the thresholds with the same level of sensitivity or specificity were also highest at the axial S1 body and lowest at L1. For example, the threshold of 90% specificity was 183 HU at the axial S1 body, 160 HU at the sagittal HU body, and 90–110 HU at L1.

Usually, we can only estimate BMD of the S1 body with the T-scores of L1–4, whereas our study provides a simple method of directly evaluating it. The HUs of the S1 body and corresponding thresholds for osteoporosis can give us more information about the bone stock of a specific S1 body before lumbar surgery. Moreover, lumbar CT is commonly performed preoperatively for patients requiring lumbar fusion; the measurement of HU demands no further investment in equipment and personnel training. Patients having high HU values can avoid extra radiation from DXA, and those having relatively lower HU values according to our thresholds can avoid being misdiagnosed with non-osteoporosis because of lumbar degeneration.

There are several limitations to our study. Firstly, it is a retrospective single-center study focusing on patients ages 50 years or older with lumbar degenerative diseases. Thus, we still need to be careful when we plan to apply the findings to other populations. Secondly, manual measurement of HUs lacks certain quality assurance protocols used in DXA and quantitative CT (QCT). Therefore, although the measurement of HU is economic and simple, it should be used as a complementary method to DXA rather than a replacement. Thirdly, the prevalence of disc degeneration at L5–S1 is higher than that at other levels, and the S1 body is relatively small compared with L1–5 vertebral bodies. As a result, the degenerative changes around or at the upper endplate of S1 might make it difficult to find appropriate ROIs of trabecular bone for measuring HUs, in which case the patients were excluded from this study.

Conclusions

Both axial and sagittal HUs of the S1 body are reliable tools with good performance in diagnosing osteoporosis in patients with lumbar degenerative diseases. Measuring HUs of the S1 body preoperatively from lumbar CT may help with surgical planning for patients with lumbar degenerative diseases.

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**Disclosures**

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

**Author Contributions**

Conception and design: Zou. Acquisition of data: Li, Zou. Analysis and interpretation of data: all authors. Drafting the article: Zou. Critically revising the article: all authors. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Li. Statistical analysis: Zou, Xu. Study supervision: Li.

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