Abstract

Objective: The aim of this study was to acquire the SUVs for each segment of the vertebral body to explore the rationale for the large variability of the SUVs in normal vertebrae.

Methods: This retrospective study was performed using the images and data from 39 cancer patients who underwent bone SPECT/CT scans with Tc-99m MDP. The SUVmax and SUVmean of the anterior, middle, posterior; left, middle, right; top, middle, and lower of the lumbar vertebrae 1~5, were calculated. The continuous variables were represented by the median values (Q1,Q3), and the differences among various segments were analyzed by Friedman’s nonparametric test. The pairwise comparison between groups was corrected by the Bonferroni method. The P<0.05 was considered statistically significant.

Results: In this study, 39 patients (31 women and 8 men; mean age, 54.79±9.12 years; age range, 39–71 years) were elucidated. The SUVmax of the L1 and L3 vertebral bodies’ parts were significantly different from those of the other portions of the same lumbar vertebra. The SUVmax of the lower posterior portion of the L1 vertebral body was significantly different from that of the upper posterior portion. There were significant differences in the SUVmax between the upper and lower portions, middle and lower portions in the middle of the L3 vertebral body. The SUVmax of the lower posterior portion of L1 and L2 vertebral were significantly different from those of the upper posterior portion. There were significant differences in the SUVmean between the upper and lower portions of the middle of the L3 vertebral body. There were significant differences among the SUVmax and SUVmean of the right, and middle of the vertebral body except for the L2 vertebral body. There were significant differences among the SUVmax and SUVmean of the anterior, middle segments of the lumbar vertebrae body, and the SUVmax of the anterior and posterior parts of the L2 vertebral body, and the SUVmax of the middle and posterior parts of the L1 vertebral body. The SUVmax of the middle and posterior portions of the L3~L5 vertebral body, and the SUVmean of the L1~L3, and L5 lumbar vertebrae had significant differences.

Conclusion: The difference in the bone metabolism of the lumbar vertebral body is caused by the variation in the horizontal direction of the vertebral body. The bone metabolism in the vertical direction of the lumbar vertebrae has more uniformity. As a quantitative imaging measure, the SUVs might require standardization with adequate reference data to minimize the variability in the participants.
### Introduction

The skeleton is the most frequent site for tumor metastasis. The extent of metastasis is the prognostic indicator in cancer patients following treatment [1,2]. Various types of tumors cause the different relative incidence of bone metastasis and the median–survival time in cancer patients [3]. The spine and pelvis are the most common sites for secondary bone metastases for different primary cancers [2,4].

The Technetium-99m labeled bone scintigraphic agent Methylene Diphosphonate (MDP) which binds to the mineral phase of bone hydroxyapatite and the calcium–rich tissue has been widely used in bone scintigraphy in metastatic bone diseases in the past [5,6]. Planar imaging has been mainly performed in bone scintigraphy, and Single-Photon Emission Computed Tomography (SPECT) has also been performed in body parts with a limited range of clinical diagnosis value. The Single-Photon Emission Tomography (SPECT)/Computed Tomography (CT) scanner combining SPECT and Computed Tomography (CT) has gained widespread acceptance. The SPECT/CT scanner provides fusion images of CT and SPECT and produces attenuation correction maps necessary for quantitative analysis using the Standardized Uptake Value (SUV) [2,5,7,8]. The SUV integrates the concept of biomarker and the notion of quantitative imaging. A Quantitative Imaging Biomarker (QIB) uses objectively measured characteristics derived from an in vivo image and serves as an indicator of normal biological processes, pathological processes, or response to a therapeutic intervention [8]. The Quantitative measurements which have been defined as the extraction of quantifiable features from the medical images to determine the normality or the degree of alteration, or the status of a disease, injury, or chronic condition relative to the normal have been vastly used with the advances in molecular imaging [8,9].

The bone scans of vertebrae were quantitatively analyzed using the SUV as QIB of SPECT/CT scans with Tc-99m-MDP [5,8,10,11]. Jun Zhao’s study evaluated SUVmax and SUVmean of benign bone lesion and malignant bone metastasis foci of normal vertebrae, which provide a quantitative reference for clinical diagnosis and the evaluation of therapeutic response in vertebral lesions [12]. The bone tissue blood flow and osteoblastic activity is proportional to bone tissue uptake of 99mTc-MDP [13,14]. bones at different sites can have different normal SUVs [5,8]. SUVmax and SUVmean were proved to be significantly different between male and female patients in most vertebrae [12]. Hence, as a quantitative imaging biomarker, SUVs of normal vertebrae showed relatively large variabilities [5,14]. Hence, as a quantitative imaging biomarker, SUVs of different between male and female patients in most vertebrae response to a therapeutic intervention [8]. The Quantitative Biomarker (QIB) uses objectively measured characteristics and the notion of quantitative imaging. A Quantitative Imaging analysis using the Standardized Uptake Value produces attenuation correction maps necessary for the SPECT/CT scanner provides fusion images of CT and SPECT. Tomography (CT) has gained widespread acceptance. The SPECT/CT scanner combining SPECT and Computed Tomography (SPECT) has also been performed in diseases in the past [5,6]. Planar imaging has been mainly used in bone scintigraphy in metastatic bone diseases [12]. The bone tissue blood flow and osteoblastic activity is proportional to bone tissue uptake of 99mTc-MDP vertebraal lesions [12]. The bone tissue blood flow and osteoblastic activity is proportional to bone tissue uptake of 99mTc-MDP.

### Materials and methods

#### Patients

In this study, we retrospectively analyzed the imaging data of patients who underwent bone scans to explore the metastasis from various cancers such as prostate cancer, pancreatic cancer, breast cancer, colon cancer, renal cancer, lung cancer, gastric cancer, and ovarian cancer in the department of Medical Image affiliated to the Hospital of the Shanxi University of Chinese Medicine. Prior to the onset of this study, the Ethics Committee of the institution approved the study protocol. Further, informed consent from patients was obtained to use the imaging data for the research purpose. Briefly, planar and SPECT/CT images were acquired about 3–4h after intravenous injection of Tc-99m–MDP. The area of the abdomen for the SPECT/CT acquisition was determined based on the patient's disease or the purpose of the examination. The data of patients with a normal lumbar vertebra who underwent bone scans were retrospectively retrieved and analyzed. Data were acquired from a group of patients undergoing Tc-99m–MDP (Guangdong Xiai nuclear pharmaceutical Co. Ltd., Xi'an, Shaanxi, China) bone SPECT/CT between August 2016 and October 2020. The patients in group were included based on the criteria according to the referred to the literature [5]: (1) access to data on measured injection activity, time of measurement, and time of injection; (2) access to patient’s weight and height information; (3) SPECT/CT that was scanned for the lumbar vertebra. The following set of exclusion criteria was outlined to determine the normal vertebrae: compression fractures, diffuse bone metastasis, ankylosing spondylitis, and metabolic bone disease. Besides, the Degenerative Joint Disease (DJD) of the spine was also characterized by correlating the tracer uptake with its CT morphological image, which encompassed osteophytes, end plates, facet joints, and the area around the joints.

#### Data acquisition and reconstruction

The Symbia T6 system (CT with a maximum of 16-slice acquisitions per rotation, Siemens Healthineers Molecular Imaging, Hoffman Estates, IL, USA) system was used to obtain SPECT/CT scans. The SPECT scans were acquired using low-energy high-resolution collimation, a 128×128 matrix of 4.8-mm pixel size, and a total of 450s/rotation in a continuous–rotation mode. After the SPECT acquisition, a low-dose CT scan was acquired with 130 kV and 15 ref mAs using adaptive dose modulation (CARE Dose 4D; Siemens Healthineers Molecular Imaging). The CT data were generated with 1.5 mm slice thickness using a smooth reconstruction B70s kernel. The SPECT reconstruction was performed using filtered back projection, and the attenuation correction was based on the attenuation maps derived from the CT data filtered by the B70s kernel.
The delineation of the Volume of Interest (VOI) was performed by a board-certified radiologist using a software “Dynamic Analysis” purchased from Beijing Dynamic Analysis Medical Information Technology Co. Ltd. (Beijing, China) which reports the statistics for the various SUVs. All the VOIs that covered the vertebral body were hand-drawn. All data were decay-corrected to the time of injection to control fluctuations of the start time of the acquisition. The final values of quantitative tracer concentrations were thus defined concerning the injection time. The patient’s clinical data were acquired from the Hospital Information System (HIS) database.

**SPECT/CT image analysis**

The Whole-Body Scan (WBS) and SPECT/CT images were independently interpreted by two experienced nuclear medicine professors and a diagnostic radiologic professor. In cases of discrepancies, the consensus was obtained by a joint reading. The SUVs of 39 lumbar vertebrae were included for the analyses based on the previously defined criteria [5]. The patients were included based on the following criteria: (1) access to data on injected activity; (2) access to patient’s weight and height information; (3) SPECT/CT scans had the lumbar vertebrae; and (4) absence of diffuse bone metastases, ankylosing spondylitis, metabolic bone disease, and osteoarthritis.

A square box with a side length of 2.71 cm (20 mL) filled with a Tc-99m MDP solution of known active concentration (approximately 2 mCi) was scanned. The data were reconstructed with filtered back-projection and the CT-based attenuation correction and then processed with SPECT/CT cameras. The SPECT/CT data were analyzed using a software “Dynamic Analysis” (Beijing Dynamic Analysis Medical Information Technology Co. Ltd., Beijing, China). The slice thickness automatically converted to be about 1.5 mm to allow isotropic voxel evaluation. A rectangular region of interest (ROI) was drawn on the cubic phantom in the central slice, measuring the SPECT count density (count/cc). The calibration factor was calculated as the ratio of actual radioactivity concentration (as measured by the dose calibrator) in the square box at the time of scanning to the measured SPECT count density per scan duration; this factor is known as the Becquerel calibration factor (BCF). The BCF is calculated as follows:

$$\text{BCF} = \frac{\text{ACC (Bq/cc) \times MC (count/cc \times 1/\text{sec})}}{\text{Injectoed activity (Bq)}}$$

The patients were scanned using the SPECT/CT system. The patient’s data in the reconstruction condition were analyzed using the same software as BCF data. The VOIs were plotted by manually adjusting the boundary of VOIs of the SPECT and the boundary of the spongy bone of the vertebral body on the CT boundaries of the fused SPECT/CT images. Then, SUVmax and SUVmean were measured [7].

**The SUV was calculated as follows:**

$$\text{SUV} = \frac{\text{BCF (Bq/cps) \times MC (count/cc \times 1/\text{Sec}) \times Body weight (g)}}{\text{Injected activity (Bq)}}$$

The vertebral body was manually divided into three parts: anterior, middle, posterior; left, middle and right; top, middle, and bottom and the SUV values of each part of the vertebral body were measured, respectively. The SPECT/CT system was calibrated with a uniform phantom, which provided a volume sensitivity factor and was specific to the camera type, collimator type, and window energy settings used. The patient’s reconstructed values were then normalized with volume sensitivity. All data were decay corrected to the time of injection to control fluctuations of the start time of the acquisition. The final values of the quantitative tracer concentrations were thus defined concerning the injection time.

**Statistical analysis**

Shapiro–Wilk test was used to determine the normal distribution of the data. The continuous variables with normal distribution were expressed as mean ± standard deviation, and one-way repeated measures of variance analyzed the differences among different groups. The continuous variables that did not follow the normal distribution were represented by the median values (Q1,Q3), and Friedman’s nonparametric test analyzed differences among various groups. The pairwise comparison between groups was corrected by the Bonferroni method. The P<0.05 was considered statistically significant. All analyses were computed using the SPSS v16.0 software (SPSS, Chicago, IL, USA). We used the Nonparametric pound–wise Multiple Comparisons module of PASS software to calculate the sample size.

**Results**

We used the Nonparametric Pair-Wise Multiple Comparisons module of PASS software to calculate the sample size. This study lumbar vertebral body SUVs between different location does not meet the normal distribution. Therefore, non-parametric tests were used to compare the differences. Assuming two comparison between three different positions, we are under the premise of type I error is less than 0.05, found 80% of the master degree of 0.5 units of the difference between any two locations, at least need to 32 cases of samples. The results are shown that 39 patients were collected retrospectively, which met the minimum sample requirements.

In this study, 39 patients (31 women and 8 men; mean age, 54.97 ± 7.69 years; age range, 39–71 years) were included from staging malignancies, such as prostate cancer, pancreatic cancer, breast cancer, lung cancer, gastric cancer, and ovarian cancer (Table 1). The lumbar vertebrae were divided into three equal parts, upper, middle, and lower parts. The SUVmax and SUVmean of the various parts of the normal lumbar vertebrae including the anterior, middle, and posterior area of the lumbar vertebra were measured based on the body weight (Figures 1-4). The normality test of all parameters showed that parameters did not obey normal distribution (P<0.001). The continuous variables that were not subject to normal distribution were expressed as median values (Q1,Q3). The SUVmax and SUVmean of the the anterior, middle, posterior; left, middle, right; top, middle, and lower, are shown in Tables 2,3. The SUVmax of the lower posterior portion of the
L1 vertebral body was significantly different from that of the upper posterior portion. In the SUVmax values between the upper and lower portions, middle and lower portions in the middle of the L3 vertebral body were significant differences. The SUVmean values of the lower posterior portion of L1 and L2 vertebral bodies were significantly different from those of the upper posterior portion. SUVmean values between the upper and lower portions of the middle of the L3 vertebral body had significant differences.

There were significant differences among SUVmax and SUVmean of the right, and middle of L1, L3, L4, and L5 vertebral bodies, but the SUVmax and SUVmean of the L2 vertebral body have no significant differences between the right and the middle portions. Significant differences were widely observed among the SUVmax of the middle and the left portions of the L1 vertebra, the SUVmean of the right and the left portions of the L1 and L4 vertebrae, and the SUVmean of the middle and left parts of the L3 vertebral body (Table 4,5). SUVmax

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<th>Table 1: Patient characteristics.</th>
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<td>Non-neoplastic patient</td>
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Discussion

The SUV might be useful as an appropriate quantitative measure in skeletal SPECT/CT imaging and has been frequently used in evaluating the activities of bone lesions.
and the response to therapy [7,15,16]. Since the SUV reflects the osteoblastic activity, and the concentration of the bone-seeking agents would be directly correlated to the SUV [13,17]. However, it is difficult to determine a standard value for a normal bone since the measurement was based on the body weight of the normal vertebral body with wide variability [5,18]. In this study, the SUVmax and SUVmean that were based on the body weight of various parts of the vertebral body such as anterior, middle, and posterior parts, and left, middle, and right segments of lumbar vertebrae were calculated.

In this current study, there were no statistical differences in most SUVs of the upper, middle, and lower parts of the lumbar vertebral body. In the human vertebrae, the trabeculae’s thickness and length of a normal cancellous bone designate the physiologic compressive and tension stresses of the bone. The trabeculae of a cancellous bone can remodel in orientation, number, and structure to accommodate and distribute the stresses to which the bone is exposed [19]. Therefore, the SUVs of the upper, middle, and lower parts of the vertebral body might have shown no statistical difference. However, the SUVmax and SUVmean of the parts of the L1 and L2 vertebral body differed significantly from those of the other portions. Since the osteoporotic fractures occur most frequently in the thoracolumbar junction (T12 to L2) due to compression, the changes in both stresses and strains were most pronounced in the posterior part of the vertebral body in this specific area [20]. Banse et al. [16] demonstrated that the vertical inhomogeneity was clearly pronounced in the images because the lower half of the vertebral body had a lower density than the upper half of the vertebral body. The differences ranged from 15% to 25% in their study. In the thoracolumbar and lumbar vertebrae, the numbers of nodes or node-to-node struts were two-fold higher in the inferior half than in the superior half of the vertebral body. The trabecular thickness and number of free-ends manifested a center or a close-to-endplate structural pattern, with the central trabeculae being 15% thicker than the close-to-endplate. Further, the central trabeculae had 30% fewer free-ends than the close-to-end plate in the same study [21]. These inferences might be the rationale for the difference in the SUVs between the upper and lower posterior segments of the L1 and L2 lumbar vertebrae. Additionally, the SUVmax of the middle portion of the L3 vertebral body had significant differences compared with the lower part and the upper part of the vertebral body. The L3 vertebral body differed in vertical inhomogeneity to a varying degree. The middle vertebral segment had more sparsely arranged trabeculae, higher porosity, and lower bone mass than the upper and lower segments of the vertebral body [22]. These characteristics of the cancellous bone of the middle segment explicate its minor role in load-bearing function. Similarly, the greatest proportion of load-bearing cortical bone appears in the smallest vertebral cross-section of the middle segment [23]. Mundinger et al. revealed that the trabeculae’s microarchitectural pattern was fine near the endplates but coarse at the center of the vertebrae [24]. Our results were consistent with the results of these previous studies relevant to the biomechanical function stating that the in vivo loads are distributed differently at the endplates compared with the middle segment of the vertebra.

This current research showed that significant differences were widely observed among SUVmax and SUVmean measures in the right, middle, and left parts of the lumbar vertebrae (Table 4). The lumbar vertebral bodies have two ossification centers, which merge and form single ossification centers on each side of the vertebral arch [25]. The TC-99m-DPD bone uptake depended on the bone osteoblastic activity, vascularization, and environmental factors [14]. In the present study, when the left, middle, and right areas of interest of the vertebral body were sketched, the vertebral body was divided into three equal parts. The ossification centers of the left and the right vertebral bodies were included in the left and right areas of interest. Therefore, the SUVs on the left and right sides of the vertebral body were greater than the SUVs on the middle part of the vertebral body. The central belt of the vertebral body is where the ossifying centers of the two sides merge. This anatomical disposition may be the reason for the significant differences among the SUVmax in the right, middle, and left portions of the lumbar vertebrae. With the advances in densitometric technologies and ex vivo analyses of the properties of a trabecular bone, it now becomes evident that the heterogeneity exists within the trabecular bone of a vertebral body [26]. The basis for the differences of the SUVmean in the right and left portions of the lumbar vertebrae may warrant further study.

The SUVmean and SUVmax measures in the middle of the vertebral body were significantly higher than those in the anterior and posterior parts of the lumbar vertebrae. There were significant differences among the SUVmean and SUVmax in the anterior, middle, and posterior portions of the lumbar vertebrae (Table 5). Banse et al. found that the structural differences between anterior, posterior, and external areas were inapparent and followed the density patterns. The cancellous bone density was 20% higher in the posterior cores than the anterior and external cores [21]. The regional morphometry and subregional density of the cancellous bones had significant differences with age and sex [26–28]. In the human vertebrae, the lumbar vertebrae play roles in both weight-bearing and load-transfer. The vertical bone trabeculae follow the lines of compressive stresses and are the primary load-bearing structure of the cancellous bone in the vertebrae, whereas the horizontal bone trabeculae are the main structure to distribute the stress and absorb the energy [19]. During aging, there is a non-uniform loss of bone within the vertebral body, resulting in non-homogeneous density throughout the centrum of the trabecular bone [19,29,30]. These sequelae may have resulted from the regional compressive load-bearing and strong energy absorption capacity of the osteoporotic population.

The main limitation of the study subjects was tumor patients without bone metastasis, which might have an inapparent selective deviation from a typically normal bone. Further, this current study had a small sample size and the changes in the SUVs due to age and sex were not considered. The study also did not compare the trend in changes in the vertebral body among radio-chemotherapy patients. The diagnostic accuracy of the absence of metastasis warrants further verification in the present study.
Conclusion

The difference in the bone metabolism of the lumbar vertebral body is caused by the variation in the horizontal direction of the vertebral body. The bone metabolism in the vertical direction of the lumbar vertebral body has more uniformity. As a quantitative imaging measure, the SUVs might require standardization with adequate reference data to minimize the variability in the participants.

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References


