Study Design: Retrospective study.

Purpose: This study aims to semiquantitatively evaluate the standardized uptake value (SUV) of $^{99m}$Tc-methylene diphosphonate (MDP) radionuclide tracer in the normal vertebrae of breast cancer patients using an integrated single-photon emission computed tomography (SPECT)/computed tomography (CT) scanner.

Overview of Literature: Molecular imaging techniques using gamma cameras and stand-alone SPECT have traditionally been utilized to evaluate metastatic bone diseases. However, these methods lack quantitative analysis capabilities, impeding accurate uptake characterization.

Methods: A total of 30 randomly selected female breast cancer patients were enrolled in this study. The SUV mean ($SUV_{\text{mean}}$) and SUV maximum ($SUV_{\text{max}}$) values for 286 normal vertebrae at the thoracic and lumbar levels were calculated based on the patients’ body weight (BW), body surface area (BSA), and lean body mass (LBM). Additionally, 106 degenerative joint disease (DJD) lesions of the spine were also characterized, and both their BW SUV$_{\text{mean}}$ and SUV$_{\text{max}}$ values were obtained. A receiver operating characteristic (ROC) curve analysis was then performed to determine the cutoff value of SUV for differentiating DJD from normal vertebrae.

Results: The mean±standard deviations for the SUV$_{\text{mean}}$ and SUV$_{\text{max}}$ in the normal vertebrae displayed a relatively wide variability: 3.92±0.27 and 6.51±0.72 for BW, 1.05±0.07 and 1.75±0.17 for BSA, and 2.70±0.19 and 4.50±0.44 for LBM, respectively. Generally, the SUV$_{\text{mean}}$ had a lower coefficient of variation than the SUV$_{\text{max}}$. For DJD, the mean±standard deviation for the BW SUV$_{\text{mean}}$ and SUV$_{\text{max}}$ was 5.26±3.24 and 7.50±4.34, respectively. Based on the ROC curve, no optimal cutoff value was found to differentiate DJD from normal vertebrae.

Conclusions: In this study, the SUV of $^{99m}$Tc-MDP was successfully determined using SPECT/CT. This research provides an approach that could potentially aid in the clinical quantification of radionuclide uptake in normal vertebrae for the management of breast cancer patients.

Keywords: Standardized uptake value; Single-photon emission computed tomography computed tomography; $^{99m}$Tc-methylene diphosphonate; Thoracic vertebrae; Lumbar vertebrae; Breast neoplasms
Introduction

The skeleton has been identified as the most frequent site for tumors to metastasize to after the treatment of primary breast cancer [1,2]. A previous study has stated that 13.6% of patients diagnosed with stages I–III breast cancer will have skeletal or bone metastases at 15-year follow-up [3]. The ribs, spine, pelvis, and upper bones of the arms and legs are among the most common sites for secondary bone metastases from primary breast cancer.

In addition to initial investigation methods such as blood cell count tests, bone scans (i.e., bone scintigraphy) can also be conducted for accurate early detection of bone metastases. A bone scan is a type of nuclear medicine-based imaging that provides an entire skeletal visualization based on the activity of a radionuclide tracer within a short amount of time. Two of the most widely used bone scan instruments are gamma cameras and single-photon emission computed tomography (SPECT), both of which are used with $^{99m}$Tc-methylene diphosphonate (MDP) tracer [4]. Conventionally, the interpretation of bone scan images has been based on a qualitative approach alone, whereby the images produced are evaluated using relative intensity values rather than absolute tracer concentration values [5]. However, the development of advanced imaging modalities, specifically the integrated SPECT/computed tomography (SPECT/CT) method, has been reported to help users obtain more information on skeletal tracer distribution due to the complementary anatomical mapping provided by the additional CT unit [6]. Moreover, the improved computing power of the SPECT/CT method has allowed for the implementation of more sophisticated image reconstruction algorithms (i.e., attenuation and scatter correction), thus enabling quantitative analysis via standardized uptake values (SUVs) to be performed [5,7].

In recent years, the introduction of SUVs into the SPECT/CT method has offered a more accessible tool for diagnostic purposes in comparison with the expensive and less available positron emission tomography (PET) method. In general, an SUV is a semiquantitative biomarker that assesses the tissue concentration of a radionuclide tracer measured by the scanner and divides it by the activity injected divided by body size [8,9]. Due to the development of the quantitative SPECT/CT technique, clinicians are now expected to be able to better differentiate metastasis and degenerative changes from the expected physiology and incidental uptake [10,11]. This could also prevent patient mismanagement [12,13]. However, regardless of the protocol published on quantitative SPECT/CT (with the earliest protocol validated on humans having been released almost 10 years ago [14]), its clinical application until now has been slow and not widely implemented [11].

Furthermore, few researchers have studied the measurement of SUV in SPECT/CT bone scans with $^{99m}$Tc tracers thus far [4-6,8,13]. Therefore, the chief purpose of this study is to report the SUV measured for the normal vertebrae of breast cancer patients undergoing $^{99m}$Tc-MDP bone scans via a SPECT/CT machine. Additionally, we also investigate the possible cutoff SUV that could be utilized to differentiate degenerative joint disease (DJD) of the spine, which constitutes benign changes, from normal vertebrae.

Materials and Methods

1. Patient selection

Patient data from a single center were retrospectively analyzed. Ethical approval, including a written informed consent waiver, was given by the institution’s Human Research Ethics Committee (USM/JEPEM/18110694) following the standards of the Declaration of Helsinki. A total of 30 female patients (mean age, 52.6±9.7 years; age range, 30–70 years) positively diagnosed with breast cancer who had undergone a $^{99m}$Tc-MDP bone scan via SPECT/CT from January 2018 to December 2018 were randomly selected based on the following inclusion criteria: (1) a SPECT/CT procedure for the thoracic spine (T1–T12) and lumbar spine levels (L1–L5) was conducted 3 hours after intravenous $^{99m}$Tc-MDP administration; (2) data available on the patient’s weight and height; (3) and data available on the injection activity and times of measurement and injection.

2. $^{99m}$Tc-MDP bone scans with SPECT/CT

The bone scans were performed using a SPECT/CT system (Discovery NM/CT 670 Pro; GE Healthcare, Chicago, IL, USA) equipped with a low-energy high-resolution (LEHR) collimator 3 hours after an intravenous injection of 736.3±22.2 MBq (19.9±0.6 mCi) $^{99m}$Tc-MDP, corresponding to 12.21±1.85 MBq (0.33±0.05 mCi/kg). A planar scan over the anterior and posterior regions of the whole body was first performed using the LEHR collima-
tor at 256×256 pixels (Fig. 1). Next, a quantitative SPECT scan with the LEHR collimator was performed at a matrix of 128×128 pixels, a step-and-shoot mode for 15 seconds per frame, and 360 rotations for every 6° view angle for the two detectors. Low-dose CT scan images were then acquired using adaptive dose modulation with 5-mm slice thickness at 120 kVp and 20 mAs.

3. Semiquantitative assessment of normal vertebrae using the standardized uptake value

The following set of exclusion criteria was first outlined to determine the normal vertebrae (i.e., healthy bone): compression fractures, diffuse bone metastasis, ankylosing spondylitis, and metabolic bone disease. In addition, DJD of the spine was also characterized by correlating the tracer uptake with its CT morphology, which encompassed osteophytes, end plates, facet joints, and the area around the joints [15]. Both the normal vertebrae and DJD of the spine were independently classified by two nuclear medicine physicians. A consensus in a joint reading resolved any discordant interpretations. The delineation of the volume of interest (VOI) was made manually using the Q.Metrix software package (GE Healthcare). Any VOIs that covered the complete vertebrae were hand-drawn from the regions of interest of three two-dimensional slices (coronal, sagittal, trans-axial), as shown in Fig. 2. SPECT images were then reconstructed using an ordered subset expectation maximization (OSEM) with depth-dependent three-dimensional (axial and trans-axial) resolution recovery, scatter correction using scatter window subtraction (triple-energy approach), and attenuation correction based on attenuation maps derived from the CT data filtered by hybrid quality control. The OSEM SPECT reconstruction used 10 subsets and two iterations without post-smoothing.

Information regarding camera sensitivity, patient data, activity injected, administration, and scan time was acquired and loaded into the Q.Metrix software before the SUV assessment. The calculated camera sensitivity was determined as 176.18 counts per minute per micro-curie (cnt/min/µCi). The SUVs were expressed as the SUV mean (SUV\textsubscript{mean}) and SUV maximum (SUV\textsubscript{max}). The SUV\textsubscript{mean} was the average multiple pixels within the VOI sampled, while the SUV\textsubscript{max} was defined as the highest pixel value within the VOI with the highest activity concentration:

\[
SUV_{\text{mean}} = \frac{\text{Total radioactivity/VOI volume}}{\text{Injected radioactivity/BW}}
\]

\[
SUV_{\text{max}} = \frac{\text{Maximum radioactivity/voxel volume}}{\text{Injected radioactivity/BW}}
\]

Note that the SUVs listed above accounted for body size or body weight (BW) measurements only. To reduce the dependence of the SUV on BW, the values for lean body mass (LBM) and body surface area (BSA) were also examined and calculated by replacing the BW parameter in both the SUV\textsubscript{max} and SUV\textsubscript{mean} [9]. Both the LBM and BSA values were automatically generated by the corresponding vendor-provided software in conjunction with the aforementioned required information.

4. Statistical analyses

All statistical analyses were performed using the IBM SPSS ver. 24.0 software (IBM Corp., Armonk, NY, USA). The degrees of dispersion of the SUV\textsubscript{mean} and SUV\textsubscript{max} for the non-rejected vertebrae (or the noise signal amount [16]) were evaluated using the coefficient of variation (CoV). The data were presented as the mean±standard deviation (SD) and analyzed using a normal distribution test. The relationship between the calculated SUVs with the patients’ age, weight, and height was evaluated using Spearman’s correlation analysis, whereby \(p\)-values of less than 0.05 indicated a significant difference. The Origin-
Pro 2018 platform (OriginLab Corp., Northampton, MA, USA) was used to graph box and whisker plots (range as a function of vertebral levels). The bottom and top of the boxes represented the first (25%) and third (75%) quartiles, respectively; the minimum and maximum values of the data were at the ends of the whiskers. The median value was marked as a line in each box, while the mean was indicated by a black circle marker inside each box. The BW SUV\textsubscript{mean} and SUV\textsubscript{max} in the treatment and no-treatment groups were also compared using an independent \(t\)-test. For DJD, the BW SUV\textsubscript{mean} and SUV\textsubscript{max} were obtained. Subsequently, the area under the curve (AUC) for both SUVs was determined from the receiver operating characteristic (ROC) curve.

**Results**

1. **SUV\textsubscript{mean} and SUV\textsubscript{max} based on BW, BSA, and LBM**

Of the 30 patients included in this study, 18 (treatment group; 60%) were identified to have undergone either a single cancer treatment or combination of treatments (chemotherapy, radiotherapy, and hormonal therapy), while the remaining 12 (no-treatment group; 40%) had...
not received any cancer treatment prior to their bone scan procedure. From the SPECT/CT scans, 286 normal vertebrae from T1 to T12 and L1 to L5 were obtained. No vertebral levels were excluded in the statistical analyses, as the sampling number was deemed adequate (n≥10). This criterion was accomplished in the same manner as the statistical considerations by Kaneta et al. [8]. The mean±SD of the BW SUV$_{\text{mean}}$ and SUV$_{\text{max}}$ was 3.92±0.72 and 6.51±0.72, respectively. Meanwhile, the BSA SUV$_{\text{mean}}$ and SUV$_{\text{max}}$ values were 1.05±0.07, and 1.75±0.17, respectively, and the LBM SUV$_{\text{mean}}$ and SUV$_{\text{max}}$ values were 2.70±0.19 and 4.50±0.44, respectively. From these results, the calculated mean values for the BW SUV$_{\text{mean}}$ and SUV$_{\text{max}}$ were found to be higher compared with those of the BSA and LBM.

The box and whisker plots of the SUV$_{\text{mean}}$ and SUV$_{\text{max}}$ for BW, BSA, and LBM at each vertebral level are shown in Figs. 3–5. The CoV values of the SUV$_{\text{mean}}$ and SUV$_{\text{max}}$ were 0.400 and 0.430 for BW, 0.390 and 0.414 for BSA, and 0.389 and 0.416 for LBM, respectively. The CoV values for each

| Vertebral level | No. | SUV$_{\text{mean}}$ | SUV$_{\text{max}}$ | SUV$_{\text{mean}}$ | SUV$_{\text{max}}$ | SUV$_{\text{mean}}$ | SUV$_{\text{max}}$ |
|----------------|-----|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|
| T1             | 13  | 0.354               | 0.391               | 0.358               | 0.388               | 0.349               | 0.385               |
| T2             | 13  | 0.376               | 0.402               | 0.380               | 0.404               | 0.358               | 0.402               |
| T3             | 16  | 0.460               | 0.488               | 0.461               | 0.501               | 0.457               | 0.501               |
| T4             | 14  | 0.495               | 0.541               | 0.480               | 0.539               | 0.468               | 0.546               |
| T5             | 15  | 0.595               | 0.599               | 0.571               | 0.583               | 0.571               | 0.591               |
| T6             | 18  | 0.479               | 0.509               | 0.462               | 0.495               | 0.462               | 0.501               |
| T7             | 20  | 0.477               | 0.472               | 0.459               | 0.451               | 0.456               | 0.453               |
| T8             | 19  | 0.452               | 0.518               | 0.436               | 0.521               | 0.437               | 0.523               |
| T9             | 18  | 0.476               | 0.535               | 0.462               | 0.529               | 0.462               | 0.534               |
| T10            | 18  | 0.443               | 0.422               | 0.434               | 0.422               | 0.434               | 0.424               |
| T11            | 20  | 0.352               | 0.339               | 0.358               | 0.326               | 0.361               | 0.329               |
| T12            | 23  | 0.354               | 0.340               | 0.359               | 0.351               | 0.357               | 0.348               |
| L1             | 26  | 0.339               | 0.351               | 0.337               | 0.365               | 0.334               | 0.360               |
| L2             | 18  | 0.306               | 0.304               | 0.359               | 0.299               | 0.307               | 0.299               |
| L3             | 14  | 0.341               | 0.341               | 0.333               | 0.321               | 0.335               | 0.324               |
| L4             | 11  | 0.247               | 0.339               | 0.236               | 0.336               | 0.242               | 0.346               |
| L5             | 10  | 0.255               | 0.408               | 0.221               | 0.203               | 0.221               | 0.205               |
| Mean CoV       |     | 0.400               | 0.430               | 0.390               | 0.414               | 0.389               | 0.416               |

CoV, coefficient of variation; SUV, standardized uptake value.
The highest CoV values of the SUV\textsubscript{mean} and SUV\textsubscript{max} among all of the vertebral levels were observed at the T5 thoracic level with 0.595 and 0.599 for BW, 0.571 and 0.583 for BSA, and 0.571 and 0.591 for LBM, respectively. Meanwhile, the lowest CoV values of the SUV\textsubscript{mean} and SUV\textsubscript{max} among all of the vertebral levels were found at the L4 lumbar level for BW (0.247, 0.339), L5 for BSA (0.221, 0.203), and L5 for LBM (0.221, 0.205).

The mean±SD of the BW SUV\textsubscript{mean} and SUV\textsubscript{max} in the no-treatment group (120 vertebrae) was 3.23±1.61 and 5.37±2.81, respectively. Comparatively, an independent t-test showed a significantly higher mean±SD ($p<0.05$) in the treatment group (166 vertebrae) with values of 4.43±1.36 and 7.36±2.49 for the SUV\textsubscript{mean} and SUV\textsubscript{max}, respectively.

2. Correlation coefficients between the SUVs and age, weight, and height

Table 2 displays the correlation coefficients between the SUVs and age, weight, and height. All SUVs were found to show a weak and no significant correlation with age. Meanwhile, there was a significant, moderate, and positive correlation between the BW SUV\textsubscript{max} with weight and a negative correlation between the BW SUV\textsubscript{mean} with height. The other SUVs indicated a weak and no significant correlation between weight and height.
3. Cutoff values of the $SUV_{\text{mean}}$ and $SUV_{\text{max}}$ in differentiating DJD of the spine and normal vertebrae

A total of 106 DJD lesions of the spine were identified. The mean±SD of the BW $SUV_{\text{mean}}$ and $SUV_{\text{max}}$ was 5.26±3.24 and 7.50±4.34, respectively. An ROC curve analysis was performed to determine the diagnostic accuracy of both SUVs in discerning DJD from normal vertebrae (Fig. 6). The AUC obtained was 0.569 (95% confidence interval [CI], 0.500–0.638) for the $SUV_{\text{mean}}$ and 0.511 (95% CI, 0.443–0.579) for the $SUV_{\text{max}}$. As the values were close to 0.5, neither SUV held a discriminatory capacity to differentiate DJD from normal vertebrae, resulting in no attainable optimal cutoff value.

Discussion

Metastases of the bone are known to not show any discernible uptake patterns during scanning [17]. The same can be said for normal bone, as the $SUV_{\text{mean}}$ and $SUV_{\text{max}}$ for each vertebral level are different and possess a considerably wide variability. A low CoV indicates a small value dispersion, while a high CoV suggests a large value dispersion. Based on the results obtained in this study, the $SUV_{\text{mean}}$ produced a lower CoV compared with the $SUV_{\text{max}}$ in all three SUV variations (BW, BSA, LBM), although the differences were rather modest. The $SUV_{\text{max}}$ is known as the standard SUV assessment method in quantitative molecular imaging (especially PET) because of its simplicity, reproducibility, resistance to partial-volume issues in tumors, and immunity to interobserver variability [9,13,18]. The $SUV_{\text{max}}$ is commonly reported in the literature, which has led to the assumption that it is the best quantitative biomarker parameter. However, it has a high dependence on the statistical quality of the images produced and, hence, is susceptible to noise [19,20]. Since CoV values illustrate the noise level in acquired data, it could be asserted that SUVs with a lower CoV (in this case, the $SUV_{\text{mean}}$) can be utilized as an optimal reference for normal bone. Recent paper of Arvola et al. [5] supports this observation, as they found higher variability in the $SUV_{\text{max}}$ values compared with those of the $SUV_{\text{mean}}$ in both $^{99m}$Tc-hydroxyethylene diphosphonate SPECT/CT and 18F sodium fluoride PET/CT scans. Therefore, despite contradicting earlier reports on the best SUV for normal bone SPECT/CT imaging [8], the $SUV_{\text{mean}}$ could also serve as an alternative in quantifying tracer uptake.

Most SUVs are measured based on BW, since this is the most popular method. In this study, the BW skeletal $SUV_{\text{mean}}$ and $SUV_{\text{max}}$ were relatively low at 3.92±0.27 and 6.51±0.72, respectively. These values were regarded as commensurable to previously reported BW SUVs for normal vertebrae, e.g., 4.4±0.5, 4.6±1.7, and 5.9±1.5 for $SUV_{\text{mean}}$, and 7.1±0.4 and 7.6±2.4 for $SUV_{\text{max}}$[8,10,21]. The $SUV_{\text{mean}}$ and $SUV_{\text{max}}$ for BSA and LBM also showed considerably low SUVs, decreasing in the order of BW, LBM, and BSA. However, none of the corrected SUVs showed a lower or higher mean CoV than those of the BW (as indicated in Table 1). Since all three SUVs yielded an almost similar result, users should select the most appropriate SUV in bone SPECT/CT imaging. If the BW SUV is used to monitor the response of $^{99m}$Tc in SPECT/CT scans, the BW should be measured with the same weighing scale at the facility rather than based on self-reported weight or patient charts [9]. The scale should also be routinely calibrated to ensure its accuracy and precision in determining patients’ BW and SUV.

In this study, based on the Spearman correlation test, it was demonstrated that all SUVs showed a weak and no significant correlation with age ($p>0.05$), similar to the results of Kaneta et al. [8]. However, Cachovan et al. [21] reported a significant negative correlation with age for all SUVs (BW and LBM) in healthy lumbar spines after $^{99m}$Tc-diphosphonate-propanedicarboxylic acid SPECT/CT scans.

### Table 2. Correlation coefficients between SUVs and age, weight, height

| Variable | Body weight | Body surface area | Lean body mass |
|----------|-------------|------------------|----------------|
|          | $SUV_{\text{mean}}$ | $SUV_{\text{max}}$ | $SUV_{\text{mean}}$ | $SUV_{\text{max}}$ | $SUV_{\text{mean}}$ | $SUV_{\text{max}}$ |
| Age      | 0.239       | 0.101            | 0.207          | 0.118          | 0.221          | 0.124         |
| Weight   | 0.158       | 0.457*           | -0.160         | 0.136          | -0.166         | -0.008        |
| Height   | -0.382*     | -0.192           | -0.315         | -0.238         | -0.299         | -0.073        |

*SUV, standardized uptake value. $p<0.05$. 

*Correlation is significant at the 0.05 level (2-tailed).
CT imaging ($p<0.001$). These variations in SUV correlation with age across many published results might be due to differences in the ages of the study populations, attenuation correction method, sample size, type of radionuclide tracer, and SPECT/CT reconstruction method. The results in this study demonstrated a significant, moderate, and positive correlation between the BW SUV$_\text{mean}$ with weight ($p<0.05$), somewhat similar to the results of Sugawara et al. [22], which found a positive correlation between BW-based blood SUV with weight upon using 2-[fluorine 18]fluoro-2-deoxy-D-glucose PET scans ($p<0.001$). This finding was likely due to the fat contribution to the BW SUV measurements [23]. Therefore, it has been suggested to use the LBM SUV to quantify the uptake of tracer in heavier or obese patients as a correction method [22].

A significant negative correlation between the BW SUV$_\text{mean}$ with height was also found in this study. Bone density composition, in general, is higher in taller subjects compared with smaller subjects. This increase of bone density has been suggested to be a result of the rise in physical burden due to the high center of gravity [8]. In a study by Axelsen et al. [24], it was concluded that chemotherapy in breast cancer patients results in a significant bone mineral densitometry (BMD) loss due to an increase in bone turnover, leading to osteoporosis. Furthermore, Huang et al. [25] also found that both the SUV$_\text{mean}$ and SUV$_\text{max}$ in osteoporotic patients are significantly lower compared with those in non-osteoporotic patients. Nonetheless, due to the retrospective design of the study, no correlations between the SUVs measured via the BMD assessment were able to be drawn. A reduction in the SUV from osteoporotic subjects might be recorded and, thus, not be representative of normal vertebrae. As such, future prospective studies incorporating BMD measurements are recommended.

Breast cancer therapy can affect normal bone homeostasis by altering the osteoclast and osteoblast functions [26]. The mechanism of action of each treatment and its effect on bone may vary, which gives rise to variation of the SUVs. In this study, the BW SUV$_\text{mean}$ and SUV$_\text{max}$ were significantly higher in the treatment group as compared with the no-treatment group. To the best of our knowledge, no study has assessed the SUVs of normal bone pre-and post-therapy; thus, further research is highly needed.

Degenerative changes of the spine, especially in the elderly, are common and often mistaken for bone metastases during bone scans. Therefore, SUV acquisition could potentially be a useful parameter to differentiate bone lesions and normal vertebrae. In a recently published paper by Mohd Rohani et al. [27] involving 34 prostate cancer patients utilizing SPECT/CT, a cutoff SUV$_\text{max}$ value of 20 was suggested to distinguish between DJD and bone metastases of the spine. However, no cutoff SUV was proposed by the authors to discriminate DJD from normal vertebrae. The results from the current study suggested that no optimal cutoff point could be identified to differentiate DJD from normal vertebrae in both the SUV$_\text{mean}$ and SUV$_\text{max}$. This finding is consistent with an earlier finding by He et al. [28] stating that there are no statistical differences between degenerative changes and normal bone in the cervical and thoracic vertebral regions.

The implementation of quantitative SPECT/CT is deemed to be more challenging compared with PET imaging due to the former’s many technical limitations. Regarding bone SPECT/CT scans, quantitative uncertainties may appear because of tracer-drug interactions (e.g., with iron supplements), metal-induced artifacts from prostheses affecting the computed SUV, variations in hormone levels such as estrogen, and unknown rates of uptake and clearance from the blood [11]. Other variables such as different radionuclide tracers with different photon energies, different collimators, and the use of different energy windows should also be considered when comparing results. The primary limitations of the current study are the small number of enrolled patients and the retrospective design, which hampered the sensitivity and specificity of the data. Moreover, SUV estimations from a completely healthy population without breast cancer were not included. Therefore, future prospective validation studies in a larger cohort are warranted to better define the SUVs of normal vertebrae.

**Conclusions**

In sum, SUVs can be used as a parameter to quantify tracer uptake in normal vertebrae SPECT/CT studies in breast cancer patients. In this research, the SUV$_\text{mean}$ and SUV$_\text{max}$ values calculated based on the patients’ BW, BSA, and LBM for normal vertebrae showed a wide variability, though they were comparable to other published results. More effort needs to be put into the clinical interpretation of SUVs to understand the quantitative nature of tracer uptake before it can be used routinely in the clinical setting.
Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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Conception and design: NMN, MFMR; data acquisition: YBU, SNBMY, MFMR, NT; analysis of data and drafting of manuscript: MFMR, SNBMY, NT; revision: NMN, WMNMZ; and supervision: NMN, WMNWZ

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