Comparison of standardized uptake values between $^{99m}$Tc-HDP SPECT/CT and $^{18}$F-NaF PET/CT in bone metastases of breast and prostate cancer

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Abstract

**Background:** Despite recent technological advances allowing for quantitative single-photon emission computed tomography (SPECT), quantitative SPECT has not been widely used in the clinical practice. The aim of this study is to evaluate the feasibility of quantitative SPECT for measuring metastatic bone uptake in breast and prostate cancer by comparing standard uptake values (SUVs) measured with $^{99m}$Tc-HDP SPECT/CT and $^{18}$F-NaF PET/CT.

**Methods:** Twenty-six breast and 27 prostate cancer patients at high risk of bone metastases underwent both $^{99m}$Tc-HDP SPECT/CT and $^{18}$F-NaF PET/CT within 14 days of each other. The SPECT and PET data were reconstructed using ordered-subset expectation-maximization algorithms achieving quantitative images. Metastatic and benign skeletal lesions visible in both data sets were identified, and their maximum, peak, and mean SUVs (SUV$_{max}$, SUV$_{peak}$, and SUV$_{mean}$) were determined. SUV ratios (SUVRs) between the lesions and adjacent normal appearing bone were also calculated. Linear regression was used to evaluate the correlations between the SUVs of SPECT and PET and Bland-Altman plots to evaluate the differences between the SUVs and SUVRs of SPECT and PET.

**Results:** A total of 231 skeletal lesions, 129 metastatic and 102 benign, were analyzed. All three SUV measures correlated very strongly between SPECT and PET ($R^2 \geq 0.80$, $p < 0.001$) when all lesions were included, and the PET SUVs were significantly higher than SPECT SUVs ($p < 0.001$). The median differences were 21%, 12%, and 19% for SUV$_{max}$, SUV$_{peak}$, and SUV$_{mean}$, respectively. On the other hand, the SUVRs were similar between SPECT and PET with median differences of 2%, −9%, and 2% for SUVR$_{max}$, SUVR$_{peak}$, and SUVR$_{mean}$, respectively.

**Conclusion:** The strong correlation between SUVs and similar SUVRs of $^{99m}$Tc-HDP SPECT/CT and $^{18}$F-NaF PET/CT demonstrate that SPECT is an applicable tool for clinical quantification of bone metabolism in osseous metastases in breast and prostate cancer patients.

**Keywords:** Quantitative SPECT, SUV SPECT, SUV comparison, SUV ratio, Bone metastases

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Background

Single-photon emission computed tomography (SPECT) has conventionally been used in a non-quantitative manner, i.e., the images have been interpreted using relative intensity values instead of absolute values of tracer concentration [1, 2]. However, the wide acceptance of integrated SPECT/CT scanners and the development of iterative reconstruction algorithms have made the clinical use of quantitative SPECT possible [3, 4]. Several applications for quantitative SPECT have been suggested but they have not been widely adopted in clinical routine use yet [1].

SPECT/CT with 99mTc-labeled diphosphonates, such as 99mTc-hydroxylamine diphosphonate (HDP), is increasingly used for the detection of bone metastases in breast and prostate cancer patients [5]. The sensitivity and specificity of SPECT/CT are superior compared to conventional bone scintigraphy or SPECT alone [6, 7]. 18F-NaF PET/CT is an even more sensitive method than 99mTc-diphosphonate SPECT/CT [6, 8], and PET is inherently quantitative [9]. However, 18F-NaF PET/CT is rather expensive and has limited availability [2, 6].

Quantitative SPECT and PET allow the calculation of standardized uptake values (SUVs), which can be utilized for longitudinal disease assessment and interpatient comparison [10]. Feasibility of quantitative bone SPECT has been previously demonstrated by reporting skeletal SUVs [2, 11–14]; however, a direct comparison to a standard reference method, such as PET/CT, has not yet been performed. 18F-NaF PET/CT has been validated using bone biopsy findings [15, 16] and proven to be repeatable in patients with bone metastases [17, 18], therefore being a potential standard method for the diagnosis of metastatic bone disease.

In this study, our aim is to explore the correlation between SUVs measured by quantitative 99mTc-HDP SPECT/CT and those measured by 18F-NaF PET/CT in the evaluation of metastatic bone uptake in breast and prostate cancer patients.

Methods

Study design and patients

This study included the same patients as a previously published study by Jambor et al. (NCT01339780, ClinicalTrials.gov) [6]. Twenty-six breast and 27 prostate cancer patients at high risk for bone metastases were prospectively enrolled between February 2011 and March 2013. Each patient gave written informed consent, and the study was approved by the local ethics committee. Patients were included if they had localized pain in the skeletal area suggesting bone metastases, suspicious laboratory findings (elevated alkaline phosphates, elevated PSA, or high PSA doubling time after prostatectomy), or suspicious histopathologic findings (stage N3a or higher in breast cancer patients, stage T3a or higher, and/or Gleason score of 4 + 3 or higher in prostate cancer patients). All patients underwent 99mTc-HDP SPECT/CT, 18F-NaF PET/CT, and 1.5-T whole-body magnetic resonance imaging (MRI), including diffusion-weighted imaging (DWI), within 14 days in a varying order. The median (range) interval between 99mTc-HDP SPECT/CT and 18F-NaF PET/CT was 3 (1–14) days [6].

99mTc-HDP SPECT/CT

The patients received intravenous injection of 672 ± 21 (mean ± SD) MBq of 99mTc-HDP. The SPECT scans were performed 180 ± 24 (mean ± SD) min after the injection using a Symbia T6, True Point SPECT/CT scanner (Siemens Healthcare, Erlangen, Germany) with the following parameters: low-energy high-resolution collimators, three bed positions, 180 projections over 360° with 9-s acquisition time per view, 128 × 128 matrix, 4.8 × 4.8 mm pixel size, and 4.8-mm slice thickness. Low-dose CT scans were acquired with 130 kV and 10 effective mAs. The CT data were reconstructed using a smooth attenuation-correction kernel B08s and a sharp bone kernel B65s.

The SPECT data were reconstructed with HybridRecon Oncology SPECT (version 1.3, HERMES Medical Solutions AB, Stockholm, Sweden) using the three-dimensional (3D) ordered-subset expectation-maximization (OSEM) algorithm with 10 iterations and 15 subsets [19]. A relatively high number of iterations was used to increase the spatial resolution of SPECT images closer to the spatial resolution of PET. The reconstruction included corrections for decay, attenuation, scatter, and collimator response. Attenuation correction was based on attenuation coefficient maps derived from the B08s CT images [20]. Scatter correction was performed with a Monte Carlo simulation using 10⁵ simulated photons and two scatter update iterations [21]. The collimator response was corrected using a Gaussian diffusion model [22]. The images were postfiltered with a Gaussian filter of 7-mm full width at half maximum (FWHM).

SPECT scanner calibration

The SPECT scanner was calibrated by scanning a uniform Jaszczak phantom (Data Spectrum Corporation, Durham, NC, USA) without any inserts inside and filled with water and 150 MBq of 99mTc-pertechnetate. A conversion factor to convert the reconstructed counts into units of activity concentration (Bq/ml) was calculated as the ratio between true activity and reconstructed counts in a homogeneous volume of interest (VOI). The resulting conversion factor was 0.107 kBq/cps.

18F-NaF PET/CT

The patients received intravenous injection of 209 ± 7 (mean ± SD) MBq of 18F-NaF. Whole-body PET scans were performed 64 ± 6 (mean ± SD) min after the injection using a Discovery VCT PET/CT or Discovery PET/CT 690 scanner (GE Healthcare, Milwaukee, WI, USA). A static emission scan from feet to skull vertex was performed with 3- (VCT
scanner) or 2-min (690 scanner) acquisition time per bed position.

The PET data were corrected for decay, attenuation, scatter, random coincidences, and dead time. The data acquired with Discovery VCT PET/CT were reconstructed in a 128 × 128 matrix with a pixel size of 5.47 × 5.47 mm and slice thickness of 3.27 mm using VUE Point HD (GE Healthcare, Milwaukee, WI, USA), a fully 3D OSEM algorithm with 2 iterations and 28 subsets. Discovery PET/CT 690 data were reconstructed in a 192 × 192 matrix with a pixel size of 3.65 × 3.65 mm and a slice thickness of 3.27 mm using VUE Point FX-Sharp IR (GE Healthcare, Milwaukee, WI, USA), also a fully 3D OSEM algorithm with 2 iterations and 24 subsets incorporating time-of-flight and point-spread function information. The images acquired with Discovery VCT PET/CT and Discovery PET/CT 690 were postfiltered with Gaussian filters of 7- and 9-mm FWHM, respectively, to lower the spatial resolution of PET closer to the spatial resolution of SPECT.

The PET scanners were quarterly calibrated following the EANM/EARL FDG-PET/CT accreditation procedure [23].

Image analysis

Image analysis was performed using HybridViewer (version 2.6, HERMES Medical Solutions AB, Stockholm, Sweden), which converted values of activity concentration into SUVs according to the following equation:

\[
SUV = \frac{\text{voxel activity concentration (Bq/ml)} \times \text{patient weight (g)}}{\text{decay corrected injected activity (Bq)} \times 1 \text{ ml/g}}.
\]  

Skeletal lesions visible in both SPECT and PET images were classified into benign and metastatic by the consensus of two experienced nuclear medicine physicians (Fig. 1). Localization CT images of PET/CT and SPECT/CT and anatomical and DWI MRI images were used to verify the lesion classification according to corresponding morphologic findings [6].

For analyzing purposes, the identified lesions were first segmented from PET images using a threshold of SUV > 15, which was lowered if the resulting VOI was clearly smaller than the area of increased uptake. Next, the same lesions were segmented from SPECT images using SUV thresholds that resulted in VOIs with the volumes similar to those in PET images.

Maximum, peak, and mean SUVs (SUV_{max}, SUV_{peak}, and SUV_{mean}) were calculated from the VOIs. SUV_{max} was determined as the SUV of the most active voxel within the VOI, SUV_{peak} as the average SUV of a 1-cm³ cube positioned within the lesion such that the enclosed average SUV was maximized, and SUV_{mean} as the average SUV of voxels in the specific VOI. The cubic VOI of

![Fig. 1](image-url)
SUVpeak could extend outside the lesion VOI in small lesions, which sometimes caused SUVpeak being lower than SUVmean.

Background bone activity was determined separately for every lesion by placing five to ten circular regions of interest (ROIs) of 1-cm diameter on adjacent normal-appearing bone tissue. These ROIs were summed to form background VOI, whose mean SUV (SUVmean,bg) was calculated. Maximum, peak, and mean SUV ratios (SUVRmax, SUVRpeak, and SUVRmean) of the lesion were calculated by dividing the SUVmax, SUVpeak, and SUVmean of the lesion by the corresponding SUVmean,bg.

Statistical analyses
Statistical analyses were performed using MATLAB (version R2016A, The MathWorks, Natick, MA, USA) and SPSS (version 24, IBM Corp., Armonk, NY, USA) software. Scatter plots and linear regression models were used to evaluate the overall correlation between SUVs of SPECT and PET data, and the Bland-Altman plots were created to evaluate the agreement between SUVs and SUVRs of SPECT and PET data. In the Bland-Altman plots, the mean difference and 95% limits of agreement (LOA) were estimated using the median and the 2.5th and 97.5th percentiles of the differences because the differences were not normally distributed according to the Shapiro-Wilk test.

The Wilcoxon signed rank test was used to determine whether the SUVs and SUVRs of SPECT and PET were statistically different, and the Mann-Whitney U test was used to determine whether the SUVs and SUVRs of metastatic lesions were significantly higher than the SUVs and SUVRs of benign lesions. These nonparametric statistical methods were employed because the SUV and SUVR data were not normally distributed. Similarly, median and interquartile range (IQR) or middle 50% were used to report SUVs.

Results
A total of 231 skeletal lesions, 129 metastatic and 102 benign, were analyzed from 46 patients. The average SUV thresholds used for segmentation were 10.7 for SPECT and 12.9 for PET. Segmented lesion volumes varied from 0.55 to 60 cm³ with a median of 2.0 cm³, and the volumes were practically the same for the corresponding SPECT and PET lesions due to the segmenta-

| Patient and lesion characteristics | Breast cancer | Prostate cancer |
|-----------------------------------|---------------|-----------------|
| Patients, n                       | 26            | 27              |
| Median age (range), years          | 61 (46–76)    | 67 (52–79)      |
| Median height (range), cm          | 167 (158–175) | 178 (169–186)   |
| Median weight (range), kg          | 70 (56–99)    | 87 (60–110)     |
| Metastatic lesions, n              |               |                 |
| 0                                 | 16            | 19              |
| 1–5                               | 4             | 6               |
| 6–10                              | 3             | 1               |
| 11–20                             | 3             | 1               |
| Benign lesions, n                  |               |                 |
| 0                                 | 5             | 2               |
| 1–5                               | 14            | 22              |
| 6–10                              | 3             | 2               |
| 11–20                             | 4             | 1               |
correlations between corresponding SUVs, but the regression lines are closer to the line of equality as their y-intercepts are almost 0. Even though the SUVR distributions of benign and metastatic lesions overlap like the SUV distributions, all SUVR measures are significantly different ($p < 0.001$) between benign and metastatic lesions.

The Bland-Altman plots for SUVR measures are shown in Fig. 5. SUVRs are even more similar than SUVs in Fig. 3 between SPECT and PET data because PET images have higher background SUVs than SPECT images (Table 3). According to the Wilcoxon signed rank test, only SUVRs peak were statistically different between SPECT and PET data. The median differences (and LOA) normalized with respect to SPECT SUVRs are 2% ($-45$–$88$%), $-9$% ($-47$–$59$%), and 2% ($-42$–$65$%) for SUVR max, SUVR peak, and SUVR mean, respectively.

**Discussion**

We evaluated the correlation and similarity of SUVs and SUVRs between $^{99m}$Tc-HDP SPECT and $^{18}$F-NaF PET in different benign skeletal lesions and bone metastases of breast and prostate cancer patients. To our knowledge, this is the first study to report SUVs and SUVRs of $^{99m}$Tc-diphosphonate SPECT using $^{18}$F-NaF PET as a reference standard. Strong correlations were shown between SUVs and similarities between SUVRs of these two methods. Previous clinical research has mainly reported $^{99m}$Tc-diphosphonate SPECT SUVs in different skeletal structures without direct reference values [2, 11–14]. Our results demonstrate that $^{99m}$Tc-HDP SPECT SUVs and SUVRs reflect the metabolic activity of lesions similarly to $^{18}$F-NaF PET SUVs and SUVRs; thus, SPECT measures can potentially be used to clinically evaluate bone metastases.
In all our analyses, PET SUVs were systematically higher than SPECT SUVs. Due to poorer spatial resolution, underestimation of SUV caused by the partial volume effect is more significant in SPECT than PET [1]. However, a higher number of OSEM iterations in SPECT and wide Gaussian filters in PET were used to compensate for this resolution difference between SPECT and PET. This compensation roughly halved the absolute differences between the SUVs of SPECT and PET. This compensation roughly halved the absolute differences between the SUVs of SPECT and PET. The compensation roughly halved the absolute differences between the SUVs of SPECT and PET. This compensation roughly halved the absolute differences between the SUVs of SPECT and PET.

After the compensation, the SUVs of PET were still significantly higher, even in background bone where no significant partial volume effect is expected. This suggests that a higher fraction of $^{18}$F-NaF than $^{99m}$Tc-HDP is extracted from blood by bone. This difference is most likely caused by the different pharmacokinetics of $^{99m}$Tc-HDP and $^{18}$F-NaF. For instance, blood protein binding hinders the extraction of $^{99m}$Tc-diphosphonates but does not affect the $^{18}$F-NaF extraction [24–26]. The SUV differences were also larger in smaller lesions, which suggests that some difference was caused by the different spatial resolutions of SPECT and PET systems even after the compensation.

In this study, SUVRs were not significantly different between SPECT and PET. Previously, the SUVR has been used mainly in the interpretation of brain PET. The SUV differences of lesions between different skeletal areas are partly caused by the differences in SUVsmean,bg between these areas, the highest SUVs and SUVsmean,bg being in the spine and the lowest in the skull. SUVs of lesions and their corresponding SUVsmean,bg cause that SUVRs are more similar than corresponding SUVs between different skeletal sites, and being also very similar between SPECT and PET. The similarity of SUVRs between SPECT and PET arises the question whether SUVRs should be used instead of SUVs for the evaluation of metastatic uptake, both in quantitative PET and especially in quantitative SPECT. SUVRs can be calculated without scanner calibration or information on the patient weight or injected dose, making them technically easier to accomplish than SUVs and equal to target-to-background ratios. However, reliable calculation of SUVsmean,bg is manually laborious and sensitive to inter-observer variability; thus, the procedure should ideally be automatized.

The typical SUV measure in PET has been SUVmax due to its simplicity and nonexistent inter-observer variability [27]. However, SUVmax is subject to noise and statistical variations in data [28]. Consequently, SUVpeak and SUVmean have gained interest as robust alternatives to SUVmax. Also in this study, SUVpeak and SUVmean were more similar than SUVmax between SPECT and PET. The strongest correlation and smallest systematic difference between SPECT and PET were in SUVpeak which is measured as an average SUV of voxels in a 1-cm$^3$ cube. Due to finite voxel size, the volumes of cubes were not exactly 1 cm$^3$. In fact, the SUVpeak of SPECT was calculated from a 26% smaller volume than the SUVpeak of PET. This somewhat compensates for the generally higher SUVs of PET, which may partly result in the smallest difference in SUVpeak. The PET SUVpeak and SUVmax as well, may have also been slightly overestimated due to collimator response correction artifact [29]. These same factors increasing SPECT SUVpeak may also cause SPECT SUVRspeak to be higher than PET SUVRspeak resulting in the largest systematic difference among our SUVR measures. However, the absolute differences we found between the SUVRs of SPECT and PET were generally very small.

### Table 2

Median (IQR) differences between SUVs of SPECT and PET for lesions in different size categories. The differences are normalized with respect to SPECT SUVs.

| Size category (cm$^3$) | n | SUV$_{max}$ (IQR) | SUV$_{peak}$ (IQR) | SUV$_{mean}$ (IQR) |
|------------------------|---|------------------|---------------------|-------------------|
| 0.35–1.21              | 58 | 41% (–22–162%)   | 17% (–20–87%)       | 30% (–12–111%)   |
| 1.32–1.98              | 58 | 24% (–24–89%)    | 11% (–39–52%)       | 19% (–34–63%)    |
| 1.98–3.97              | 58 | 17% (–45–106%)   | 7% (–45–76%)        | 17% (–39–90%)    |
| 3.97–60.31             | 57 | 11% (–29–83%)    | 6% (–26–52%)        | 17% (–14–51%)    |

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### Table 3

SUVsmean,bg from normal appearing bone at different skeletal sites.

| Skeletal site | PET SUV median (IQR) | SPECT SUV median (IQR) | PET SUV–SPECT SUV median (IQR) |
|--------------|----------------------|------------------------|--------------------------------|
| Skull        | 2.6 (2.1–5.0)        | 2.0 (1.8–2.6)          | 0.3 (0.1–0.8) (p < 0.001)      |
| Spine        | 7.4 (6.6–8.7)        | 6.1 (4.7–6.8)          | 1.7 (0.6–2.8) (p < 0.001)      |
| Rib cage     | 3.0 (2.4–4.0)        | 2.5 (2.0–3.1)          | 0.7 (0.1–1.0) (p < 0.001)      |
| Pelvis       | 5.5 (4.7–6.6)        | 4.7 (4.1–6.0)          | 0.6 (0.2–1.3) (p < 0.01)       |
| Limbs        | 3.8 (2.9–4.9)        | 3.2 (2.1–4.3)          | 0.7 (0.1–1.0) (p < 0.01)       |

IQR interquartile range

$^p$ value denotes at which level a median difference is significantly different from 0 according to the Wilcoxon signed rank test.
The strong correlation of SUVs and very similar SUVRs between $^{99m}$Tc-HDP SPECT and $^{18}$F-NaF PET suggest that the SUVs and SUVRs of $^{99m}$Tc-HDP SPECT are valid and feasible for research and clinical use. These results encourage SPECT SUVs to be utilized in clinical trials and follow-up studies to decrease inter-observer variability and to standardize SPECT results between patients, imaging systems, and clinical centers. Even though $^{18}$F-NaF PET/CT is a more sensitive method than $^{99m}$Tc-HDP SPECT/CT, the cost of $^{18}$F-NaF PET/CT is three- to four-fold that of $^{99m}$Tc-HDP SPECT/CT [6], making $^{99m}$Tc-HDP SPECT/CT the more cost-effective modality for the evaluation of metastatic bone uptake in breast and prostate cancer. SPECT has also wider availability than PET in most countries [2]. The SUV regression lines have slopes so close to one that the SUVs of SPECT and PET would even be equal in some cases if a small positive bias was added into SPECT SUV. However, because the Bland-Altman plots show rather wide 95% LOA, PET SUVs cannot be directly converted into SPECT SUVs. Therefore, usage of SPECT as follow-up for PET is not recommended.

The requirements for producing quantitative SPECT and PET data are the same. The most important ones are scanner calibration and a reconstruction algorithm correcting for photon attenuation and scatter within the object [1]. The calculation of reliable SUVs also requires accurate measurements of prepared activity, preparation time, injection time, syringe residual activity, time of residual activity measurement, and patient weight.

The main limitations of our study are the relatively small number of patients with bone metastases and the lack of accurate injected activities used in SPECT. We have afterwards measured the distribution of injected...
activities from a sample of 38 syringes used for bone scans at our department. Our average injected activity should be 670 MBq. In that estimation, the injected activity was calculated as the activity of full syringe subtracted by the residual activity, and it had an average of 672 MBq and SD of 21 MBq. This average injected activity was eventually used in the SPECT SUV calculations of this study. We assume that the inaccuracy in the assessment of injected activity results in a maximum random error of 7% (= 2 SDs) in the individual SPECT SUV calculations and no error in the SPECT SUVR calculations. This error in SPECT SUV calculations is relatively small compared to SUV differences between SPECT and PET, so it does not have a significant influence on the statistical differences between SPECT and PET SUVs.

Conclusion
The SUVs of $^{99m}$Tc-HDP SPECT/CT and $^{18}$F-NaF PET/CT correlate strongly and SUVRs are very similar, showing that SPECT SUVs are feasible for uptake measurements in bone metastases of breast and prostate cancer. SUVs and SUVRs could be used to expand the visual analysis of skeletal SPECT especially in follow-up studies. The SUVs of $^{99m}$Tc-HDP SPECT/CT are slightly lower than the SUVs of $^{18}$F-NaF PET/CT.

Abbreviations
3D: Three-dimensional; CT: Computed tomography; DWI: Diffusion-weighted imaging; FWHM: Full width at half maximum; HDP: Hydroxyethylidiphosphonate; IQR: Interquartile range; LOA: Limits of agreement; MRI: Magnetic resonance imaging; DSWM: Ordered-subset expectation-maximization; PET: Positron emission tomography; PSA: Prostate-specific antigen; ROI: Region of interest; SD: Standard deviation; SPECT: Single-photon emission computed tomography; SUV: Standardized uptake value; SUVR: Standard uptake value ratio; VOI: Volume of interest

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Availability of data and materials
The data supporting the findings of this study are included in this published article.

Authors’ contributions
SA and TN designed the study. All authors participated in the data acquisition. SA and IJ analyzed the data. SA and TN drafted the manuscript. IJ, AK, JK, SK, and MS commented on the manuscript, and all authors approved the final version of the manuscript.

Ethics approval and consent to participate
This study was approved by the Ethics Committee of the Hospital District of Southwest Finland and each patient gave written informed consent.

Consent for publication
Each patient gave written informed consent for publication.

Competing interests
The authors declare that they have no competing interests.

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