The influence of using different reconstruction algorithms on sensitivity of quantitative $^{18}$F-FDG-PET volumetric measures to background activity variation

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ABSTRACT

Introduction: This study aims to investigate the influence of background activity variation on image quantification in differently reconstructed PET/CT images.

Methods: Measurements were performed on a Discovery-690 PET/CT scanner using a custom-built NEMA-like phantom. A background activity level of 5.3 and 2.6 kBq/ml $^{18}$F-FDG were applied. Images were reconstructed employing four different reconstruction algorithms: HD (OSEM with no PSF or TOF), PSF only, TOF only, and TOFPSF, with Gaussian filters of 3 and 6.4 mm in FWHM. $S_{\text{UV}_{\text{max}}}$ and $S_{\text{UV}_{\text{peak}}}$ were obtained and used as cut-off thresholding; Metabolic Tumor Volume (MTV) and Total Lesion Glycolysis (TLG) were measured. The volume recovery coefficients (VRCs), the relative percent error ($\Delta$MTV), and Dice similarity coefficient were assessed with respect to true values.

Results: $S_{\text{UV}_{\text{max}}}$ and $S_{\text{UV}_{\text{peak}}}$ decreased and MTV increased as function of increasing the background dose. The most differences occur in smaller volumes with 3-mm filter; Non-TOF and Non-PSF reconstruction methods were more sensitive to increasing the background activity in the smaller and larger volumes, respectively. The TLG values were affected in the small lesions (decrease up to 12%). In a range of target volumes, differences between the mean $\Delta$MTV in the high and low background dose varied from -11.8% to 7.2% using $S_{\text{UV}_{\text{max}}}$ and from 2.1% to 7.6% using $S_{\text{UV}_{\text{peak}}}$ inter reconstruction methods.

Conclusion: The effect of the background activity variation on SUV-based quantification in small lesion was more noticeable than large lesion. The HD and TOFPSF algorithms had the lowest and the highest sensitivity to background activity, respectively.

Key words: $^{18}$F-FDG-PET/CT; Background activity; Image reconstruction; MTV; TLG

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INTRODUCTION

Positron emission tomography (PET) based molecular imaging, as a tool with the potential to improve diagnosis of cancer, determining tumor volume and localization, treatment planning and/or treatment response assessment, plays an increasingly valuable role in radiation oncology [1-3]. During the past two decades, the glucose analogue $^{18}$F-Fluorodeoxyglucose (FDG) as an appropriate radiotracer was applied. In the most studies, the maximum standardized uptake value ($SUV_{max}$) is the most widely used index for tissue accumulation of $^{18}$F-FDG to quantification of normal and tumor tissues [1]. Peak SUV ($SUV_{peak}$) has been suggested as a more robust alternative to $SUV_{max}$ in cancer studies reports, as well [1]. The use of SUVs is easy, but these rely on a single (or few) voxels and do not reflect the metabolic information of the whole tumor volume. Unlike SUVs, Metabolic tumor volume (MTV) depicts the tumor biological target volume [4]. MTV is a SUV-based parameter which defined as the volume of tumor tissues consists of the total number of voxels that have uptake above a predetermined SUV threshold. Recently, studies have consistently reported MTV can provide valuable prognostic information, and may better reflect the overall tumor burden for various neoplasms from surrounding normal tissues compared with SUVs and conventional anatomic approaches [1, 5-8]. MTV does not indicate the clonogenic cells density within the tumor volume; Total Lesion Glycolysis (TLG) is another quantitative volumetric metrics that combines the volumetric and metabolic information defined as MTV multiplied with the average SUV ($SUV_{mean}$ which is an index of the clonogenic cells density) [9, 10].

Recently, it has been demonstrated that combined PET/computed tomography (CT) imaging is feasible and useful for PET/CT-guided radiation therapy planning [2, 3, 11]. Advanced treatment techniques such as PET-based intensity-modulated radiation therapy, allow additional dose escalation (i.e. dose-painting by intensity-modulated radiation therapy) to improve tumor control probability without increasing organ at risks toxicity but rely on reliable tumor volume delineation based on accurate molecular imaging [3, 12-14]. PET/CT imaging-based dose painting is a concept to prescription and delivery of non-uniform radiation dose which radiation dose is shaped according to the PET uptake [15, 16]. Also, the extraction and characterization of various volumetric textural features (Radiomics) with analysis of PET/CT images was suggested for tumor response assessment [17, 18]. Therefore, such applications relied on an accurate segmentation and reliable delineation of the tumor volume and treatment response assessment [19-22].

At present, the use of fixed thresholding-based methods of the SUVs (especially $SUV_{max}$) is more common in clinical FDG-PET/CT imaging and hold a significant role in biological target delineation and assessing patient response to cancer therapy [23, 24]. PET/CT based biological target delineation remains highly variable and it is still subject to debate [25, 26]. It is known that PET/CT segmentation and reconstruction algorithms can lead to significant differences in measuring MTV and accurate tumor delineation [25, 27]. One that can affect the extracted biological target volume from both segmentation and reconstruction algorithms is the additional background FDG uptake by the tumor surrounding tissues, which can affect SUV-based parameter and lead to significant noise, contrast loss and some uncertainties in tumor burden detection and accurate tumor delineation [28-30].

The impact of several factors such as the tumor size, the image reconstruction methods used for image generation or the tumor to background ratio has been evaluated in a number of studies on oncological patients [25, 31-33]. Background or surrounding normal tissues activity uptake in patients is unavoidable and can vary greatly between organs [29], and it should be taken into account during PET/CT based target volume delineation [34]. In particular there are limited publications about the impact of background activity of the surrounding normal tissues on volumetric parameters when using different reconstruction methods, moreover, typical evaluation approaches in the assessment of volumetric accuracy or overlap measures (such as volume recovery coefficient, dice similarity coefficient, and etc.) were proposed [35].

In the current work, we used the $^{18}$F-FDG-PET images of a custom-built NEMA-like phantom to determine the influence of background activity variation on quantification of SUV and variability of SUV-based volumetric parameters (including MTV and TLG) in the PET images reconstructed using different reconstruction parameters. Both $SUV_{max}$ and $SUV_{peak}$ are applied as image indices for thresholding.

METHODS

Data acquisition and image reconstruction

FDG-PET/CT imaging was performed using a PET/CT Discovery 690 (GE Healthcare, Milwaukee, USA) includes a 64-slice CT scanner. The baseline fully 3D ordered subset expectation maximization (3D-OSEM) algorithm (referred to as HD), system-specific point-spared-function (PSF), and time-of-flight (TOF) information are provided by the manufacturer. FDG-PET raw data were reconstructed with four different reconstruction algorithms: i) HD = without TOF and PSF, ii) PSF = HD + PSF, iii) TOF = HD + TOF, and iv) TOFPSF = HD + TOF + PSF.
The standard 3D iterative HD algorithm with post-smoothing Gaussian filter 3 and 6.4 mm in the full width at half maximum (FWHM) was applied (HD3 and HD6.4, respectively). In addition, PSF, TOF, and TOFPSSF which incorporates the PSF and/or TOF within the standard HD algorithm were used with post-smoothing Gaussian filter 3 and 6.4 mm FWHM (PSF3, PSF6.4; TOF3, TOF6.4; TOFPSSF3, TOFPSSF6.4, respectively). In total, 8 different PET/CT reconstructions methods with the same raw data were available. HD6.4 was applied as the default reconstruction method with 3 iterations and 18 subsets. For all datasets, TOF reconstructions were performed with 2 iterations and 18 subsets and non-TOF reconstructions were performed with 3 iterations and 18 subsets. 10 min for each bed position was used in all reconstruction methods. The coincidence time window was 4.1 ns, and the TOF time resolution was 555 ps. The image matrix was 256 x 256; in-plane voxel size was always 2.73 x 2.73 mm (with slice thickness 3.27 mm). The CT data with 120 kV and 100 mA were acquired for attenuation correction.

**Phantom studies**

**Imaging protocol:** A custom-made phantom, resembling the National Electrical Manufacturers Association (NEMA) image quality phantom with six fillable cylinders was used for all measurements. The cylinders with 25 mm height and diameter of 10, 13, 17, 22, 28 and 37 mm were initially filled with a solution of $^{18}$F-FDG with the background activity level of 5.3 kBq / ml and a tumor-to-background ratio of 4:1. To determine the impact of low-dose background activity, the activity decay time was subsequently calculated to reach the background activity level of 2.6 kBq / ml. Laser markers were used for the phantom positioning in the center of the PET/CT scanner field of view. Measurements were performed over two bed positions in 3-dimensional list mode with 10 min per a bed scan.

**Data analysis:** The SUV normalized to body mass were calculated for all reconstruction methods. Later, the SUV$_{\text{max}}$ and the SUV$_{\text{peak}}$ were calculated: The SUV$_{\text{max}}$ was a voxel with the highest intensity in each target volume; SUV$_{\text{peak}}$ was calculated using a 3-dimensional 1.0 ml spherical VOI which was positioned on a high-uptake region within the tumor such that the average value is maximized [36].

MTV was calculated by thresholding-based image segmentation methods with the 50% cutoff thresholding value for all reconstruction at two background activity level. This cutoff thresholding value was chosen as the threshold ratio to achieve MTV closest to the actual volume [36]. MTV was extracted from the voxels inscribed with the VOI on the condition SUV $\geq$ 50% $\times$ SUV$_{\text{max}}$.

The size of the VOI was considered the same as the physical shape of cylinders in the phantom. The actual cylinders volumes were measured and with respect to the true volume, volumetric accuracy in each specific tumor size was calculated using the volume recovery coefficient (VRC), defined as the following equation:

$$VRC = \frac{MTV_{\text{recon},i}}{\text{True Volume}_j} \quad \text{Eq. 1}$$

Here, MTV$_{\text{recon},i}$ refers to the MTV corresponding to the $i^{th}$ reconstruction method; True Volume$_j$ refers to the actual size of $j^{th}$ cylinder volume. The actual volumes for each cylinder correspond to its known physical volume were 2, 3.5, 5.5, 10.5, 16, and 28.5 cm$^3$ for volume with the diameter of 10, 13, 17, 22, 28 and 37 mm, respectively.

The relative percent error in volume (called $\Delta$MTV) with respect to true volume was calculated as follows:

$$\Delta$MTV = \frac{MTV_{\text{recon},i} - \text{True Volume}_j}{\text{True Volume}_j} \times 10 \quad \text{Eq. 2}$$

In addition, the overlap between reconstructions and real target volumes were calculated by Dice’s similarity coefficient (DSC) [37]:

$$\text{DSC} = \frac{2(MTV_{\text{recon},i} \cap \text{True Volume}_j)}{MTV_{\text{recon},i} + \text{True Volume}_j} \times 100 \quad \text{Eq. 3}$$

**Target size-dependent comparison:** In order to compare the inter-method variability at two background activity level with considering the tumor sizes, MTV and subsequently VRC, $\Delta$MTV, and Dice’s similarity coefficient were calculated. Later, the quantitative value of TLG was obtained to assess the impact of background activity variation on clonogenic cells density calculation. TLG was defined as the MTV multiplied with the average SUV (SUV$_{\text{mean}}$) in the MTV.

**Target size-independent comparison:** To assess the influence of background activity variation on MTV for a range of target sizes (i.e. without considering the tumor sizes), the mean percent error $\Delta$MTV and standard deviation (SD) for each reconstruction method was calculated in the range of our phantom target sizes. SD was used as a criterion to assess the volume dependency of each reconstruction method.

All calculations were repeated and compared on the condition SUV $\geq$ 50% $\times$ SUV$_{\text{peak}}$ for all reconstruction methods and target volume size at two background activity level. In some cases, the results for the smallest (10 mm cylinder diameter), medium (17 mm cylinder diameter), and largest (37 mm cylinder diameter) volumes are presented.
RESULTS

Figure 1a illustrates the $SUV_{max}$ and $SUV_{peak}$ as a function of cylinder diameter for different reconstruction methods at two background activity levels (Low Dose: 2.6 kBq / ml; High Dose: 5.3 kBq / ml). The results showed by increasing the background activity level, the relative difference of SUVs had different behaviors whenever the target volume diameter was less than about four times the FWHM of the imaging system resolution. So, the mean relative difference of SUVs in the cylinder with diameter ≤ 17-mm and diameter >17 mm is shown in Figure 1b.

Figure 2a shows the VRCs for each cylinder diameter and reconstruction methods using the cut-off thresholds $SUV_{max}$ and $SUV_{peak}$ at two background activity levels. The results on the mean relative difference of VRCs are compared in Figure 2b for the cylinder with diameter ≤ 17-mm and diameter > 17 mm. The result showed that the VRC for small target volumes was more sensitive to increasing background activity than large target volumes.

A summary of percent error $\Delta MTV$ (%), as relative differences in real target volume and MTV, in the low and high background activity for different target volumes and reconstruction methods are shown in Figure 3; the smallest (10 mm diameter), the medium (17 mm diameter), and the largest (37 mm diameter) volumes are presented with cut-off 50% of $SUV_{max}$ and $SUV_{peak}$.

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Fig 1. a) $SUV_{max}$ and $SUV_{peak}$ as function of cylinder diameter (D) for different reconstruction methods at two background activity levels (Low Dose: 2.6 kBq / mL; High Dose: 5.3 kBq / mL). Sources-to-background ratio of 4:1 was used. b) The mean relative difference of SUVs by increasing the background activity level in the cylinder with D ≤ 17-mm and D > 17-mm. SUV relative difference (%) = ($SUV_{High Dose}$ - $SUV_{Low Dose}$) / $SUV_{Low Dose}$ * 100

Figures and captions are included in the provided text to illustrate the results and methods used in the study.
Fig 2. a) Volume recovery coefficients (VRC) as function of cylinder diameter (D) for different reconstruction methods at two background activity levels (Low Dose: 2.6 kBq/mL; High Dose: 5.3 kBq/mL). b) the mean relative difference of VRC by increasing the background activity level in the cylinder with $D \leq 17$ mm and $D > 17$ mm. VRC relative difference (%) = $\frac{(VRC_{High\ Dose} - VRC_{Low\ Dose})}{VRC_{Low\ Dose}} \times 100$

The overlap between each true target volume and MTV of reconstruction methods were calculated by Dice’s similarity coefficient; Figure 4 details the data on Dice similarity coefficient for each specific target volume at two background activity levels.

TLG was calculated in each target volume to analyze the target size-dependent variation in different background activity levels; Table 1 summarizes these data on the smallest, medium and largest volumes for different reconstruction methods. The relative difference of TLG in two background activity levels as biases (%) is presented in Table 1. The maximum bias occurred when 3-mm filter and $SUV_{max}$ was used for all four algorithms.

Table 2 compares mean percent error $\Delta$MTV and corresponding mean Dice similarity coefficient in a range of target volumes for eight reconstruction methods and two different background activity levels. Also, inter-method dependencies to target volume size as SD are compared in Table 2.

**DISCUSSION**

We have analyzed the influence of background activity variation on quantification of SUVs (i.e. $SUV_{max}$ and $SUV_{peak}$) and variability of SUV-based volumetric parameters (i.e. MTV and TLG) in different PET reconstruction methods.
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Fig 3. Percent error ΔMTV (%) for each specific tumor size for different reconstruction methods at two background activity levels (Low Dose: 2.6 kBq / mL; High Dose: 5.3 kBq / mL). The smallest (10 mm diameter), the medium (17 mm diameter), and the largest (37 mm diameter) volumes are presented; a) $\text{SUV}_{\text{max}}$, b) $\text{SUV}_{\text{peak}}$

Fig 4. Dice similarity coefficient for each specific tumor size for different reconstruction methods at two background activity levels (Low Dose: 2.6 kBq / mL; High Dose: 5.3 kBq / mL). The smallest (10 mm diameter), the medium (17 mm diameter), and the largest (37 mm diameter) volumes are presented; a) $\text{SUV}_{\text{max}}$, b) $\text{SUV}_{\text{peak}}$
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Table 1: TLG measurements for different reconstruction methods at two background activity levels (Low Dose: 2.6 kBq/mL; High Dose: 5.3 kBq/mL). The bias (%) represents the relative difference of TLG in two background activity levels. Cut-off 50% of SUV$_{\text{max}}$ (upper half) and cut-off 50% of SUV$_{\text{peak}}$ (lower half).

<table>
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<td>10</td>
<td>17</td>
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<tr>
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<td>110.8</td>
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TLG = MTV × SUVmean; Bias (%) = ((TLG$_{\text{low Dose}}$ – TLG$_{\text{high Dose}}$)/ TLG$_{\text{low Dose}}$) × 100; Diameter (volume): 10 mm (2 cm$^3$), 17 mm (5.5 cm$^3$), 37 mm (28.5 cm$^3$)

Although, previous studies have demonstrated the image reconstruction and segmentation methods can affect the PET image quantification and make it difficult to detect tumor burden [25, 33, 38], but it is necessary to know the inter-reconstruction method variability of image quantification at different background (i.e. surrounding normal tissues) activity level, and it should also be taken into account during PET/CT based target volume delineation. Our results showed that even though the variability in SUVs and SUV-based volumetric quantification were small, the background activity level can affect target volume quantification in some reconstruction method.

SUV$_{\text{max}}$ and SUV$_{\text{peak}}$ decreased as function of increasing background activity level. Figure 1a indicates exactly where SUV$_{\text{max}}$ and SUV$_{\text{peak}}$ meet the reference SUV (i.e. SUV = 4 calculated based on decay corrected tracer activities) within target volumes. There was a clear trend in the mean relative difference of SUVs, as shown in Figure 1b. Non-TOF reconstruction methods (i.e. HD and PSF) were more sensitive to increasing the background dose in the cylinder with diameter ≤ 17-mm. Also, there was a greater sensitivity to increasing the background dose in 17-mm < diameter for Non-PSF reconstruction methods (i.e. HD and TOF). The number of events increases by increasing the activity of the tumor surrounding tissues; as well as, the random and scatter events increase. This effect typically causing the noise to be distributed along the line of response, and it also reduces the SUV. As shown in Figure 1a, SUV$_{\text{max}}$ in HD reconstruction algorithm with the small filter was most susceptible to an increase in background activity. The TOF include reconstruction method with localizes the decay site based on the arrival time of the photons at the detector provides adequate temporal resolution. The PSF algorithm by incorporating more information about the detector system response (i.e. detector geometry, parallax effects and all of which contribute to a spatially variant blurring point response) improves spatial resolution and improves contrast recovery. So, TOF and PSF along with HD algorithm compensating the SUV reduction whether they were in combination or separated. SUV$_{\text{max}}$ in TOF and TOFPSF reconstruction methods less affected by noise and it had more stability. Variability of SUV$_{\text{max}}$ was more than SUV$_{\text{peak}}$ which was due to less statistical variation and noise effects in SUV$_{\text{peak}}$.

Table 1: TLG measurements for different reconstruction methods at two background activity levels (Low Dose: 2.6 kBq/mL; High Dose: 5.3 kBq/mL). The bias (%) represents the relative difference of TLG in two background activity levels. Cut-off 50% of SUV$_{\text{max}}$ (upper half) and cut-off 50% of SUV$_{\text{peak}}$ (lower half).
Table 2: Mean percent error ΔMTV (%) and corresponding Dice similarity coefficient on all target volumes in different background activity level and reconstruction methods. Cut-off 50% of SUV$_{\text{max}}$ (upper half) and cut-off 50% of SUV$_{\text{peak}}$ (lower half).

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<td>38.4 151.9 -6.3 63.1</td>
<td>0.86 0.57 0.99 0.17</td>
<td>46 182.1 -5.8 74.2</td>
<td>0.84 0.52 0.99 0.19</td>
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<tr>
<td>TOF3</td>
<td>17.4 92 -6.6 38.3</td>
<td>0.91 0.68 0.97 0.11</td>
<td>22 104.8 -6 43.3</td>
<td>0.9 0.66 0.98 0.13</td>
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<tr>
<td>TOF6.4</td>
<td>37.4 148.2 -4.1 59.2</td>
<td>0.87 0.57 0.99 0.17</td>
<td>41.1 152.4 -2.5 60.9</td>
<td>0.86 0.57 0.99 0.17</td>
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<tr>
<td>TOFPSF3</td>
<td>14.1 94.3 -13.5 41.6</td>
<td>0.9 0.68 0.99 0.11</td>
<td>18.8 107.6 -15.5 46.7</td>
<td>0.89 0.65 0.98 0.12</td>
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<tr>
<td>TOFPSF6.4</td>
<td>33.3 142.9 -7.6 58.9</td>
<td>0.87 0.58 0.99 0.16</td>
<td>35.3 148.6 -7.2 61</td>
<td>0.86 0.57 0.98 0.16</td>
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</table>

SD: Standard Deviation; ΔMTV: The relative percent error in metabolic tumor volume
In small lesions, the small filter size was more suitable and the behavior of reconstruction methods was important. For all four reconstruction algorithms, the VRCs plots illustrated that the VRCs were close to 1.0 using $\text{SUV}_{\text{max}}$ with 3-mm filter size in both background activity levels, although at a higher background activity was slightly closer. With increasing background activity, the lowest and the largest change in VRCs were seen in TOFPSF and HD reconstruction methods in the small volumes using $\text{SUV}_{\text{max}}$ cut-off threshold, respectively. Reconstruction with 6.4-mm filter size less affected by the background activity variation and lower inter-reconstruction methods variability observed (Figure 1b) which is due to averaging and suppressing the noise effect. However, in small lesions, as shown in Figure 2, the use of larger post-smoothing filter can lead to lesion volume overestimation (1< VRCs), so that VRCs in different reconstruction algorithms obtained up to 1.6 - 1.9 in low, and 1.7 - 2 in high background activity levels. The reconstruction with 6.4-mm filter size not suitable for small tumor quantification, although they showed less sensitivity to increase the background dose (Figures 1 and 2).

For all reconstruction methods, VRCs was severely increased in diameter ≤ 17-mm using $\text{SUV}_{\text{peak}}$ cut-off thresholds. These increases were up to 1.9 - 2.6 in low, and 2 - 2.8 in high background activity levels. The $\text{SUV}_{\text{peak}}$ cut-off threshold was not suitable for small lesions (diameter ≤ 17-mm) quantification. However, inter-reconstruction methods variability in the cylinder with diameter > 17 mm was negligible.

The value of MTV was increased by increasing the background activity level. This increase was significant in small target volumes. In 10-mm diameter cylinder using $\text{SUV}_{\text{max}}$, MTV increases in percent for HD3, PSF3, TOF3, and TOFPSF3 were 23.1%, 20.6%, 17.4%, and 12.7%, respectively; and for HD6.4, PSF6.4, TOF6.4, and TOFPSF6.4 were 5.6%, 4.6%, 11.3%, and 9.3%, respectively. By increasing the background activity, the percent error ΔMTV was reduced or increased when ΔMTV was negative or positive, respectively (equation 2 and Figure 3).

A high degree of overlap was observed between true target volumes and corresponding MTVs at different background dose. As can be seen in Figure 4, by increasing the background dose, Dice similarity coefficient has slightly increased or decreased depending on whether the primary MTV was larger or smaller than true target volume. The largest changes in the Dice similarity coefficient were observed in a smaller volume when $\text{SUV}_{\text{max}}$ was used as the cut-off threshold. In the smallest target volume, Dice similarity coefficients were 0.88, 0.97, 0.87, 0.94 in the low background dose and 0.98, 0.94, 0.95, 1.00 in the high background dose for HD3, PSF3, TOF3, and TOFPSF3, respectively. For cutoff using $\text{SUV}_{\text{max}}$, the use of TOFPSF algorithm with a small post-smoothing filter in the presence of a background activity uptake can improve the Dice similarity coefficient. The large filter in small tumor volumes leads to a reduction in the Dice similarity coefficient, whether the $\text{SUV}_{\text{max}}$ was cutoff value or $\text{SUV}_{\text{peak}}$. Using the large filter in the presence of background activity uptake lead to more uncertainty in the tumor quantification and/or delineation.

As detailed in Table 1, the TLG value for large target volume was not significantly changed by increasing background activity level. Increasing the background activity from 2.6 kBq / ml to 5.3 kBq / ml in the smallest target volumes resulted in the largest TLG difference in HD3, PSF3, TOF3, and TOFPSF3 with -12.0%, -10.1%, -7.5%, and -4.0% bias (=100 × (TLG low Dose − TLG high Dose) / TLG low Dose). Hence, the quantification of TLG was also affected by reconstruction methods and target volume.

In a range of target volumes, as can be seen in Table 2, the differences between the mean ΔMTV in the high and low background dose (i.e. mean ΔMTV high Dose - mean ΔMTV low Dose) varied from -11.8% to 7.2% using $\text{SUV}_{\text{max}}$ and from 2.1% to 7.6% using $\text{SUV}_{\text{peak}}$ inter reconstruction methods. The mean ΔMTV at two background dose showed small variations within one type of reconstruction method (i.e. TOFPSF); however, there was a significant difference in the mean ΔMTV in different reconstruction methods.

With increasing the surrounding background uptake for all four algorithms, the SD (i.e. tumor size dependency of each reconstruction method) increased on average about 20.6% for filter 3-mm and 16.0% for filter 6.4-mm using $\text{SUV}_{\text{max}}$, and 12.1% for filter 3-mm and 8.6% for filter 6.4-mm using $\text{SUV}_{\text{peak}}$, respectively. PSF3 with 30.7% and TOF3 with 27.2% increases in SD had the maximum variation by increasing background dose using $\text{SUV}_{\text{max}}$.

There are many PET segmentation methods for delineation of radiation therapy treatment volumes which some of them use the surrounding background activity uptake and analyzed the impact of varying signal-to-background ratios on tumor quantifications [39-41]. One study investigated the relation between variation in background intensity on PET-based gross tumor volume delineation and SUVs in normal lung, aorta (blood pool), and liver tissues [29]. Those investigators also came to the conclusion that it is reasonable and precise to tumor volume delineation in patients after taking into account the background intensity of the lung lobe in which the tumor is found. This conclusion agrees with our results showing that a variation in background activity can cause vast variation in the SUVs and the SUV-based volumetric measurements. In addition, our results showed some reconstruction methods were more sensitive to
background dose especially in small lesions as described in detail. The problems of segmentation approaches in tumor volume delineation in FDG-PET have addressed in the literature [24]. Other investigators have looked at the other tumor segmentation requirements, but the results are controversial and they did not take into account the impact of different reconstruction algorithms and post smoothing filters size [24]. One of the most important limitations of thresholding approaches for PET-image segmentation is that does not consider background activity. However, despite many other methods, thresholding approaches are more common in clinics and they have an intuitive basis. So, the authors suggest that fixed thresholding-based segmentation methods require considering the reconstruction method and post smoothing filters size.

There are some limitations to this study. Our phantom study was performed in a range of simple shape volumes (2-28 cm$^3$) and finding might not be transferable to smaller volumes or lesion with a complex shape in clinical situations. Only one tumor-to-background ratio was considered, the background activity uptake in low contrast lesion may be most problematic.

CONCLUSION

Our study addresses the questions of volumetric tumor quantification and/or delineation in different background activity uptake and different reconstruction methods both for SUV$_{\text{max}}$ and SUV$_{\text{peak}}$ cut-off thresholds. The effect of the background activity variation on SUV-based quantification in small lesion was more noticeable than large lesion. SUV$_{\text{max}}$ and a smaller filter were more suitable whenever the tumor size was less than three to four times the FWHM of the reconstructed image resolution. The HD and TOFPSP algorithms had the lowest and the highest sensitivity to background activity variation, respectively.

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