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Effect of Time-of-Flight and Regularized Reconstructions on Quantitative Measurements and Qualitative Assessments in Newly Diagnosed Prostate Cancer With $^{18}$F-Fluorocholine Dual Time Point PET/MRI

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Abstract
Recent technical advances in positron emission tomography/magnetic resonance imaging (PET/MRI) technology allow much improved time-of-flight (TOF) and regularized iterative PET reconstruction (RIR) algorithms. We evaluated the effect of TOF and RIR on standardized uptake values (maximum and peak SUV [SUV$_{max}$ and SUV$_{peak}$]) and their metabolic tumor volume dependencies and visual image quality for $^{18}$F-fluorocholine PET/MRI in patients with newly diagnosed prostate cancer. Fourteen patients were administered with 3 MBq/kg of $^{18}$F-fluorocholine and scanned dynamically for 30 minutes. Positron emission tomography images were divided to early and late time points (1-6 minutes summed and 7-30 minutes summed). The values of the different SUVs were documented for dominant PET-avid lesions, and metabolic tumor volume was estimated using a 50% isocountour and SUV threshold of 2.5. Image quality was assessed via visual acuity scoring (VAS). We found that incorporation of TOF or RIR increased lesion SUVs. The lesion to background ratio was not improved by TOF reconstruction, while RIR improved the lesion to background ratio significantly ($P < .05$). The values of the different VAS were all significantly higher ($P < .05$) for RIR images over TOF, RIR over non-TOF, and TOF over non-TOF. In conclusion, our data indicate that TOF or RIR should be incorporated into current protocols when available.

Keywords
cancer detection imaging, novel imaging methods/agents for clinical studies, quantitation in molecular imaging, cancer imaging, PET/MRI, prostate cancer

Introduction
Prostate cancer remains the most common cancer diagnosed in men in the United States and is among the leading causes of cancer-related mortality.¹ Imaging continues to play an increasing role in the evaluation of prostate cancer and suspected cancer recurrence. Multiparametric prostate magnetic resonance imaging (MRI) is the current diagnostic imaging workhorse, although it is still undergoing continuous evolution to overcome inherent limitations. For example, Muller et al recently showed that the revised Prostate Imaging Reporting and Data System (PI-RADS 2.0) provides moderately reproducible MRI scores similar to PI-RADS 1.0 for clinically relevant prostate cancer.² Newer experimental radiopharmaceuticals have made positron emission tomography (PET) combined

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with computed tomography (CT) and MRI interesting research subjects aimed at improving the diagnostic accuracy of prostate cancer diagnosis.3-5

Combined with multiparametric prostate MRI, PET imaging in PET/MRI is a natural extension of current prostate imaging practice.6,7 Recent advancements in PET detector technology also have made time-of-flight (TOF) imaging capability and its associated TOF-enabled reconstruction, which is a standard technology in modern PET/CT scanners, feasible in the clinical PET/MRI setting. The TOF-enabled reconstruction (hereinafter, “TOF reconstruction” for simplicity) is known for improved spatial resolution, contrast to noise ratio, and image quality in PET/CT.8-13 The improved resolution for TOF reconstruction, in comparison to that for non-TOF reconstruction (ie, the reconstruction without the TOF feature), is primarily due to faster convergence; thus at comparable iterations, TOF reconstruction typically yields better spatial resolution than non-TOF reconstruction does.

Further technological advance in TOF-PET, particularly the use of solid-state photomultipliers instead of photomultiplier tubes as the photodetector, made it possible for the TOF capability to be implemented even in a strong magnetic field, translating it into clinical PET/MRI systems as well.12,14-16

In addition to TOF reconstruction capability, a regularized iterative reconstruction (RIR) algorithm has been clinically implemented (Q.Clear; GE Healthcare, Waukesha, Wisconsin).17-20 However, this regularized reconstruction algorithm has not been available clinically for PET/MRI yet. Currently, no literature exists examining the effect of TOF or regularized reconstruction on standardized uptake values (SUVs), image quality, or lesion conspicuity for PET/MRI imaging of prostate cancer. Also, there is no standard static acquisition protocol for the clinically used prostate cancer PET radiopharmaceuticals such as 11C-acetate, 18F-fluorocholine, and 68Ga-labelled Glu-urea-Lys(Ahx)-HBED-CC (68Ga–HBED-CC).3-5,21 We acquired our 18F-fluorocholine PET/MRI dynamically over 30 minutes,18F-fluorocholine uptake was recorded in list mode acquisition -flex MR images (repetition time [TR] = 4 millisecond, echo time [TE]: 2.33 millisecond, flip angle: 5° or 12°, partial Fourier: 70.3%, acquisition time: 18 seconds) with a standard body coil developed for PET/MRI.

(ie, CAPRA score ≥ 3) were evaluated with 18F-fluorocholine PET/MRI of the pelvis in this institutional review board–approved study. Dominant clinically relevant tumors with Gleason score 3+4/4+3 or greater were evaluated. Prostatic lesions were found in 12 patients, and a total of 17 tumors were identified by 18F-fluorocholine PET/MRI for quantitative analysis.

18F-Fluorocholine PET/MRI Protocol

Positron emission tomography imaging was performed immediately following intravenous (IV) administration of 3 MBq/kg of 18F-fluorocholine via a peripheral IV catheter after at least 4 hours of fasting for 30 minutes. Positron emission tomography images were acquired on a TOF-PET/3-tesla MRI scanner (SIGNA PET/MR; GE Healthcare) in 3-D acquisition mode over the pelvis. 18F-fluorocholine uptake was recorded in list mode and the list-mode data were replayed into 2 time points, 1 to 6 minutes summed and 7 to 30 minutes summed after administration of the radiotracer. Based on the observation of the tracer kinetics over 30 minutes from dynamic reconstructions of the entire 30 minutes,16 the early time duration (1-6 minutes), excluding the first minute of mostly blood pool uptake, was determined since the rapid uptake of 18F-fluorocholine reaches a plateau mostly around 5 to 6 minutes. In addition, the late time duration (7-30 minutes), with at least a minute gap from the early time point, was determined to capture accumulated activity after the radiotracer reached the plateau. Reconstruction was performed in a matrix size of 128 × 128 (voxel size = 2.34 × 2.34 × 2.78 mm3) with other parameters: transverse field of view (TR) = 300 mm, TOF-enabled and TOF-disabled OS-EM algorithm with 28 subsets and 2 iterations, 5-mm full-width at half-maximum postreconstruction Gaussian filter and 1:4:1 axial filter for both TOF-enabled reconstruction (“TOF reconstruction”), and TOF-disabled reconstruction (“non-TOF reconstruction”). Finally, the RIR algorithm was used for the same data sets with a regularization parameter beta value of 350, 3 and 2 initial non-TOF regularized and OS-EM iterations followed by 8 TOF regularized iterations. A set of representative images using non-TOF, TOF, and RIR algorithms for both time points (early and late) are shown in Figure 1. The choice of reconstruction parameters such as the number of iterations, the number of subsets, and the sequence of non-TOF and TOF regularized iterations were based on the vendor-default recommended parameters which are commonly used at our center as well, not intended to be optimized further by our own investigation. This way, we assessed the image qualities from each reconstruction algorithm as all of the readers using this imaging system would see. Magnetic resonance imaging was performed simultaneously during PET acquisition utilizing an endorectal coil. Standard MR-based attenuation correction was applied using liver accelerated volume acquisition -flex MR images (repetition time [TR] ~ 4 millisecond, echo time [TE]: 2.23 millisecond, flip angle: 5° or 12°, partial Fourier: 70.3%, acquisition time: 18 seconds) with a standard body coil developed for PET/MRI.
Maximum and peak SUVs were documented for dominant suspicious prostate lesions and SUVmean was obtained of the pelvis (ischium) as background. All SUVs were measured on Osirix (Pixmeo, Bernex, Switzerland). Metabolic tumor volume by PET was measured using a scale of 1 to 100. The paired 2-tailed t test was used to compare the SUV ratios, TBRs, TBloods, and VAS’s to investigate whether the difference is statistically significant.

### Results

#### Positron Emission Tomography Image Analyses

All reconstructed PET images were reviewed independently by 2 board-certified radiologists with training in nuclear medicine. Metabolic tumor volume by PET was measured using a 50% isocountour and an SUV threshold of 2.5 to determine tumor boundaries and calculate the encompassed volume utilizing Osirix. For quantitative analysis, we first examined the difference in SUV measurements from different reconstruction for early (1-6 minutes) and late time (7-30 minutes) point data by taking the ratios of lesion SUVmax and SUVpeak from TOF and RIR reconstructions to those from non-TOF reconstructions.

Since an RIR reconstruction with a sufficiently large number of iterations results in a higher spatial resolution than corresponding TOF and non-TOF reconstructions, and a TOF reconstruction also converges faster than non-TOF reconstruction with the same number of iterations for both, the relative SUVs are expected to depend on the size of lesion where SUVs were derived. It is important to note that for RIR reconstruction, the number of iterations should be sufficiently large as in our case (3 and 2 initial non-TOF regularized and OS-EM iterations followed by 8 TOF regularized iterations) because RIR reconstruction can provide poorly converged images (ie, smoother images) accompanied by spatial resolution degradation. Hence, we also examined the lesion volume dependencies of these ratios by generating scatter plots of the ratios over the metabolic tumor volumes.

Then, we compared the lesion to background (target to background ratio or TBR) and lesion to blood ratios (target to blood or TBlood) from different reconstructions and different time point data to illustrate any perceived visual contrast of lesion to background with regard to different reconstruction and different time points. After this step, in order to depict how different reconstructions and different time points affect TBR and TBlood, we took ratios of TBRs and TBlood of TOF to those of non-TOF and ratios of TBRs and TBlood of RIR to those of non-TOF reconstructions for both time point (early and late) data sets. Finally, image quality was subjectively assessed using visual acuity scoring (VAS), ranking image quality on a scale of 1 to 100. The paired 2-tailed t test was used to compare the SUV ratios, TBRs, TBloods, and VAS’s to investigate whether the difference is statistically significant.

#### Standardized Uptake Values Consistency

Relative lesion SUVs for TOF and RIR data sets to non-TOF data sets (SUV_TOF/SUV_non-TOF, SUV_RIR/SUV_non-TOF) are as follows. The ratios of SUVmax for TOF to that for non-TOF were 1.14 ± 0.29 (range: 1.00-1.72) for the 1- to 6-minute uptake period and 1.17 ± 0.13 (range: 1.02-1.45) for the 7- to 30-minute uptake period, respectively. The ratios of SUVmax for RIR to that for non-TOF were 1.56 ± 0.42 (range: 1.20-2.64) for the 1- to 6-minute uptake period and 1.26 ± 0.42 (range: 1.00-2.43) for the 7- to 30-minute uptake period, respectively. The ratios of SUVpeak for TOF to that for non-TOF were 1.14 ± 0.13 (range: 0.98-1.49) for the 1- to 6-minute uptake period and 1.11 ± 0.10 (range: 0.93-1.30) for the 7- to 30-minute uptake period, respectively. The ratios of SUVpeak for RIR to that for non-TOF were 1.30 ± 0.21 (range: 0.97-1.96) for the 1- to 6-minute uptake period and 1.06 ± 0.20 (range: 0.70-1.52) for the 7- to 30-minute uptake period, respectively. Figure 2 shows the SUV ratios for all reconstructions and at the 2 time points in 1 plot to depict the relative consistencies of SUV measurements. The relative lesion SUVs of RIR to non-TOF reconstructions are significantly larger than those of TOF to non-TOF in the early time point images (P < .05); however, the relative lesion SUVs of RIR to non-TOF reconstruction do not show statistically significant differences to those of TOF to non-TOF reconstruction in the late time point images (P > .05).

#### Standardized Uptake Values Dependence on Metabolic Volume

Using the metabolic volume average from 2 methods (50% isocountour and 2.5 SUV threshold) measured for the data sets reconstructed with TOF and RIR, scatter plots were generated to show the ratios of lesion SUVs (SUVmax and SUVpeak) of TOF to non-TOF and RIR to non-TOF reconstructions,
respectively (Figure 3). These scatter plots clearly show the dependence on the measured volume for the relative SUVs. The smaller the metabolic tumor volume is, the larger relative SUVs were observed for both TOF and RIR reconstructions over non-TOF reconstructions. In addition, for the RIR reconstructions as they are expected to have better convergences than both TOF and non-TOF reconstructions, the difference in SUVs is greater in small metabolic volumes. These data also show that when the metabolic tumor size reaches approximately 4 mL, the relative SUVs do not show increase over the volume.

Lesion to Background and Lesion to Blood

Relative TBRs for TOF and RIR data sets to non-TOF data sets ($\text{TBR}_{\text{TOF}} / \text{TBR}_{\text{non-TOF}}$, $\text{TBR}_{\text{RIR}} / \text{TBR}_{\text{non-TOF}}$) and relative TBloods for TOF and RIR data sets to non-TOF data sets ($\text{TBlood}_{\text{TOF}} / \text{TBlood}_{\text{non-TOF}}$, $\text{TBlood}_{\text{RIR}} / \text{TBlood}_{\text{non-TOF}}$) are as follows. The ratios of TBR for TOF to that for non-TOF were $0.86 \pm 0.25$ (range: 0.59-1.47) for the 1- to 6-minute uptake period and $0.85 \pm 0.13$ (range: 0.54-1.02) for the 7- to 30-minute uptake period, respectively. The ratios of TBR for RIR to that for non-TOF were $1.03 \pm 0.35$ (range: 0.59-1.89) for the 1- to 6-minute uptake period and $1.04 \pm 0.31$ (range: 0.64-1.58) for the 7- to 30-minute uptake period, respectively. The ratios of TBlood for TOF to that for non-TOF were $0.96 \pm 0.14$ (range: 0.75-1.19) for the 1- to 6-minute uptake period and $1.07 \pm 0.12$ (range: 0.83-1.20) for the 7- to 30-minute uptake period, respectively. The ratios of TBlood for RIR to that for non-TOF were $1.53 \pm 0.35$ (range: 0.78-2.37) for the 1- to 6-minute uptake period and $1.20 \pm 0.33$ (range: 0.77-2.01) for the 7- to 30-minute uptake period, respectively. Figure 4 shows the TBRs and TBloods for all reconstructions and at the 2 time points in 1 plot to depict the changes and consistencies of lesion to background and lesion to blood contrast. The relative TBRs and TBloods of RIR over non-TOF are all larger than those of TOF over non-TOF reconstructions at all time points. The differences are all statistically significant ($P < .05$) except for the difference between TBlood of RIR over non-TOF and TBlood of TOF over non-TOF for the 7- to 30-minute data set ($P > .05$).

Image Quality

Mean VAS was 57.6, 68.1, and 80.9 for non-TOF, TOF, and RIR reconstructions, respectively for one of the radiologists,
and 58.8, 70.9, and 78.6, respectively, for the other radiologist for the 1- to 6-minute acquisition data sets and 48.4, 53.4, and 76.5, respectively, for one of the radiologists and 55.1, 68.4, and 75.4, respectively, for the other radiologist for the 7- to 30-minute acquisition data sets. The VAS was significantly higher for RIR reconstructed data sets over TOF reconstructed data sets, TOF over non-TOF, and RIR over TOF for both time point data sets (P < .05). The VAS was better for the early time point data (1-6 minutes summed) than the late time point data (7-30 minutes summed) for non-TOF reconstructions with statistical significance (P < .05) for both readers; however, for TOF and RIR reconstructions, there was no statistically significant difference (P > .05) between the 2 time point data. Figure 5 shows box plots of VAS between data sets for the 2 radiologists who performed VAS.

**Discussion**

Addition of TOF and RIR capability for image reconstruction to 18F-fluorocholine PET/MRI increases SUV\textsubscript{max} and SUV\textsubscript{peak} for both 1- to 6-minute and 7- to 30-minute acquisition data sets over corresponding non-TOF data sets. The reason for this difference is primarily because of improved spatial resolution of TOF-enabled PET reconstruction when the same number of iterations was used in our case and improved spatial resolution of RIR PET reconstruction when a sufficient number of iterations were performed. Consistent with a modest increase in SUV\textsubscript{max} and SUV\textsubscript{peak} using TOF and RIR reconstructions, image quality assessed by VAS also showed significant improvement when TOF and regularized reconstructions were used over images generated by conventional non-TOF reconstructions. Some of these findings are in keeping with the previously reported benefits of TOF for PET/CT, including increased signal and contrast to noise ratio, improved lesion detectability, and detecting low contrast lesions in a noisy background by increasing sensitivity and reducing image noise.\textsuperscript{9,25-27} These benefits of TOF should allow for a decrease in patient dose, acquisition time, or both as demonstrated following the addition of TOF to PET/CT.\textsuperscript{28} Unlike extensive literature reports on the benefits of TOF reconstructions, reports of regularized reconstruction’s benefits in quantitative measurements and qualitative visual image assessment using clinical data are scarce; however, our findings are consistent with a few case reports in the literature.\textsuperscript{19,29}

There are several limitations inherent to this study, including small sample size and retrospective nature. Also, our assessment was only confined to prostate 18F-fluorocholine PET/MRI data sets, while the TOF and RIR algorithms can be used for any other PET data sets when these algorithms are available. Effects of TOF and RIR on clinical outcomes and sensitivity of detecting intermediate or high-grade prostate cancer were not assessed. While radiotracer uptake was noted in all

![Figure 4](image1.png)

**Figure 4.** Target-to-background (TBR, lesion-to-ischium) and target-to-blood (TBlood, lesion-to-blood pool) differences are shown as ratios of TBRs and TBloods of TOF data sets to non-TOF data sets and RIR data sets to non-TOF data sets for both early and late time point images. RIR indicates regularized iterative reconstruction; TOF, time-of-flight.

![Figure 5](image2.png)

**Figure 5.** Box plots of VAS performed by 2 board-certified radiologists. VAS indicates visual acuity scoring.
lesions, uptake pattern in nontarget lesions, such as benign prostatic hypertrophy nodules, was not assessed. It is possible that the increase in $SUV_{\text{max}}$ and $SUV_{\text{peak}}$ with TOF and RIR could translate into an increased sensitivity in detecting small lesions, though this topic requires further investigation.

**Conclusion**

Our data strongly suggest that incorporation of TOF and RIR algorithms should be used when available for prostate $^{18}$F-fluorocholine PET/MRI. In particular, RIR algorithm outperformed TOF algorithm without regularization when compared side-by-side in terms of image quality assessment for our data sets. Hence, in order to fully capture the promise of RIR algorithms, further investigation on the performance of RIR including TOF for other PET imaging scenarios is warranted.

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**Declaration of Conflicting Interests**

The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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