Feasibility assessment of yttrium-90 liver radioembolization imaging using amplitude-based gated PET/CT

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Introduction
Liver cancer is estimated to cause over 23,000 deaths per year in the USA and is expected to grow by 2.9–6.5% in the next 15 years [1]. Radioembolization using yttrium-90 (90Y) spheres is a globally expanding palliative treatment option for patients with hepatocellular carcinoma [2]. SPECT imaging is typically used for assessment [3]; however, current systems lack robust quantitative capabilities, inhibiting their ability to produce accurate dosimetry calculations [4]. 90Y-PET/computed tomography (CT) imaging following radioembolization has gained momentum as a method for dose assessment because of its ability to provide accurate quantitative dosimetry [5–7], despite low count rates from the low frequency of internal pair production [8] and image degradation caused by normal breathing patterns [9]. PET/CT dosimetry estimates can be used to quantitatively assess whether a given therapy delivery was successful or not [10].

Traditional phased-based gating can correct for motion, but results in noisier images with counts divided across each gate. Amplitude-based gating, available commercially as HD-Chest (Siemens Medical Solutions USA Inc., Knoxville, Tennessee, USA), is a software package that enables the ability to select a percentage of the amplitude of the respiratory cycle measured by a respiratory band that determines the data used in the final reconstruction [11]. Data reconstructed using amplitude gating methods have reduced motion compared with static acquisitions (Fig. 1), but with 2–5-fold less count loss compared with traditional phase-based gating [12]. This method has been shown to be effective in reducing motion artifacts and improving quantitative accuracy in routine clinical PET imaging [13]. In this technical note, we show the initial results that suggest that this technique may improve quantitative measurements without sacrificing potential diagnostic integrity because of increased image noise.

Patients and methods

Patient population
Informed consent was obtained from ten patients and the study was carried out under the auspices of a University of Tennessee Graduate School of Medicine Institutional Review Board-approved study (#3502). Patients ranged in age from 60 to 80 years and all underwent 90Y radioembolization for the treatment of primary or metastatic hepatocellular carcinoma, followed by 90Y PET/CT imaging. All patients in the study received SirSpheres
radioembolization therapy (SIRTeX Medical Limited, North Sydney, Australia) with standard of care lung shunt determinations [14] used to assess that they clinically qualify for radioembolization therapy.

Dose calculations
Dose-deposition estimates were calculated manually using the local deposition model. This model can be used to scale the activity concentration values measured using PET imaging to estimates of $^{90}$Y dose deposition using the following equation:

\[
D_{90Y} (\text{Gy}) = A_0 \left( \frac{\text{Bq}}{\text{ml}} \right) \times K_{90Y} \left( \frac{\text{Gy} \times \text{ml}}{\text{Bq}} \right),
\]

where the conversion factor $K$ for $^{90}$Y has been found previously in other research to be $4.782 \times 10^{-5}$ Gy/ml/Bq [7]. Using these methods, an estimate was obtained for the $^{90}$Y dose deposited for each patient following their $^{90}$Y radioembolization procedure.

Imaging protocol
All patients were imaged on a 64-slice, 4-ring Biograph mCT Flow (Siemens Medical Solutions USA Inc.). A 20–45 min listmode acquisition over the liver was performed for each patient using continuous bed motion acquisition. An Anzai respiratory band (Anzai Medical, Tokyo, Japan) was fitted around each patient to measure the respiratory cycle. The band was placed $\sim 5$ cm from the top of the navel for each patient.

PET data were histogrammed into static (standard) and amplitude-based gated (corrected) datasets. Corrected data were reprocessed using the scanner default 30–35% amplitude threshold for reconstruction. All PET data were decay, branching ratio, attenuation, and scatter corrected using the standard systems and software provided on the Biograph mCT platform. No additional manual corrections were applied that could potentially introduce other biases not associated with the standard scanner hardware and software. Data were reconstructed using resolution recovery algorithms and time-of-flight corrections (200×200 image matrix, 1 iteration, 21 subsets, 2 mm Gaussian filter per previous assessments performed on this system [7]). Nongated CT data were acquired for anatomical segmentation at 120 kVp using a continuously modulated tube current and 5 mm slices (512×512 image matrix, 1 mm$^3$ Gaussian filter).

Data analysis
Three methods were used to assess the impact of amplitude gating on delivered $^{90}$Y dose deposition estimates compared with static imaging. Two of the assessments use PET data voxel value thresholding to draw region of interests (ROIs) using CT segmented liver regions as threshold boundaries. The third assessment
used CT data to segment individual primary lesions and use those ROIs transferred to the PET image voxels to assess 90Y dose deposition. All regions of interest were drawn using the Inveon Research Workplace (Siemens Healthineers, Knoxville, Tennessee, USA).

Two threshold settings were used for image analysis as represented in Fig. 2. The first method used a standard style image threshold that is used in commonly nuclear medicine analysis with a lower bound determined by 30% of the maximum value in the lesion (henceforth referred to as the 30 maximum threshold method). The second method (region-based threshold method) used a lower bound derived from the maximum of several ROIs drawn in the healthy parenchyma of the embolized liver, with the upper bound of both thresholds set to the maximum pixel value in the reconstructed PET image. This method enables improved comparison of static and gated data as increases in high-frequency noise will also propagate into the regions used to set the lower threshold limit. The third method (individual lesion method) involved segmentation of 17 individual primary lesions across all patients using the fused PET/CT data and voxel values from the PET images used for conversion to 90Y dose deposition estimates. For each ROI, the mean, minimum, maximum, and SD of the activity concentration were recorded as well as the ROI volume. ROI activity concentration values were then multiplied by the above-mentioned conversion factor of 4.78 × 10^{-5} to convert from activity concentration (Bq/ml) into an absorbed dose estimate (Gy) [15]. Measured values were compared statistically between static and amplitude-gated scan types as increases in high-frequency noise will also propagate into the regions used to set the lower threshold limit. The second method (region-based threshold method) used a lower bound derived from the maximum of several ROIs drawn in the healthy parenchyma of the embolized liver, with the upper bound of both thresholds set to the maximum pixel value in the reconstructed PET image. This method enables improved comparison of static and gated data as increases in high-frequency noise will also propagate into the regions used to set the lower threshold limit. The third method (individual lesion method) involved segmentation of 17 individual primary lesions across all patients using the fused PET/CT data and voxel values from the PET images used for conversion to 90Y dose deposition estimates. For each ROI, the mean, minimum, maximum, and SD of the activity concentration were recorded as well as the ROI volume. ROI activity concentration values were then multiplied by the above-mentioned conversion factor of 4.78 × 10^{-5} to convert from activity concentration (Bq/ml) into an absorbed dose estimate (Gy) [15]. Measured values were compared statistically between static and amplitude-gated scan types as well as the final recorded estimates of the 90Y dose obtain from the final therapy radiology reports.

Diagnostic assessment
Data were assessed clinically by a board-certified nuclear medicine physician using the methods described by Kao et al. [10], which are diagnostic criteria developed to assess the quality of the 90Y therapy delivery as opposed to the diagnosis of the patient’s disease. A binary scoring system based on Kao et al.’s [10] methods, shown in Table 1, was created with the inclusion of an additional parameter to compare activity ‘bleeding’ on standard and corrected datasets. A score of 1 indicated that the given criteria were fulfilled, whereas a score of 0 indicates that the criteria were not fulfilled. Cumulative scores were compared between methods to assess any changes in diagnostic image interpretation. A window/level threshold of 25% of maximal intensity was used for all patients for the visual diagnostic assessment.

Results
Quantitative assessment
ROI volumes were found to be significantly reduced for all threshold methods when using motion-corrected data versus the standard images. Volumes of ROIs drawn on motion-corrected images were reduced by an average of 36 and 79% percent for 30 maximum and region-based threshold methods, respectively, in the global liver analyses. Individual lesion analysis yielded average volume reductions of 16%. The maximum reductions observed for global and local lesion analysis were greater than 40 and 60%, respectively. Statistics collected for volume changes are shown in Table 2.

Mean dose calculations in global liver analyses were shown to be underestimated by an average of 23% using region-based thresholding (Table 3). Individual lesion analysis indicated an average underestimation of dose of 8%. Higher standard deviations were observed in corrected images, with the average increase for region-based threshold methods being 12% for global liver calculations and 15% for individual lesion assessments. Calculations of coefficient of variation indicated a reduction of 19 and 28%, respectively.

With one exception, all measurements of maximum values comparing standard images with those using amplitude-based gating were statistically significant with P value less than 0.05. Post-hoc power analysis indicated

Table 1 Binary rating scale on the basis of the Kao criteria

| 90Y Binary criteria | 0/1 |
|---------------------|-----|
| Technical success criteria fulfilled – 0/1 |
| Unsuccessful criteria not fulfilled – 0/1 |
| Nontarget activity criteria not fulfilled – 0/1 |
| Noise spike criteria not fulfilled – 0/1 |
| Uncorrected motion ‘bleeding’ – 0/1 |

The Kao criteria are a clinical assessment of diagnostic data acquired from postradioembolization PET/CT. For this work, we developed a binary scale on the basis of these criteria to enable a quantitative assessment of the diagnostic quality of the therapy delivery.

CT, computed tomography; 90Y, yttrium-90.
that we attained more than 90% and more than 80% power for global and local assessments, respectively, even with the small number of patients used in this initial assessment. Values of estimated maximum dose are presented in Table 4. All percent changes in values are presented in Table 5.

The mean and standard deviation of estimated delivered doses recorded in the radiology reports for all 10 patients following radioembolization was 130.74 ± 56 Gy. The 30 maximum threshold method resulted in underestimation of the actual dose for both static and amplitude-gated datasets with underestimations of 37 and 34%, respectively. For the region-based thresholding method, static data underestimated dose by 20% whereas amplitude-gated data underestimated dose by only 5%. These percentages can be derived from the mean calculated dose values found in Table 3.

### Diagnostic assessment

Diagnostic assessments of data indicated an observed increase in image noise for motion-corrected data, with noise spike criteria fulfilled in 100% of all motion-corrected cases compared with 18% for standard static imaging. Criteria for technical success, according to the Kao criteria, were fulfilled in all patients analyzed, with no patients showing signs of unsuccessful treatment or off-target activity. Bleeding of activity into areas around treatment regions in nonmotion-corrected PET images was observed in 82% of cases. Figure 3 shows a comparison image that shows the reduced blur in motion-corrected images at the expense of additional noise.

### Discussion

Quantitative metrics show that the amplitude gating methods described here generally increased values for dose estimates from PET/CT imaging, which may be underestimated because of respiratory motion. Although image noise increases as a result of count loss when using gating methods, the coefficient of variation decreased by an average of 19%. This suggests that improved event localization led to higher mean values that helped compensate for increased noise. ROI volumes also showed statistically significant reductions for threshold-based methods, which is most likely the result of the combined effects of both decreased motion blur making apparent tumor volumes smaller and the impact of increased high-frequency noise in the data from reduced counts.

Diagnostic scoring of images showed that the increase in image noise in corrected images did not result in a change in the determination of whether a procedure was successful or unsuccessful. Also, increased noise did not have any effect on the assessment of off-target activity, which could be a concern with radioembolization procedures [16].

Statistical significance was observed when comparing standard and motion-corrected values, with an estimated post-hoc power of 0.9 and 0.8 for global and individual lesion measurements, respectively. One measurement in the global analysis was observed to be nonsignificant, but was determined to likely be the result of a type II statistical error.

Standard imaging showed an average underestimation of maximum dose of 40%, which may be significant if using 90Y PET/CT to verify 90Y dose. This work shows initial results that suggest that the use of amplitude-based respiratory motion correction may be a useful tool for post 90Y-radioembolization PET/CT.

### Conclusion

The use of amplitude-based gated PET/CT imaging in post-90Y radioembolization assessment is clinically feasible. We have shown initial results that amplitude-based gating can be used to improve dose estimates with minimal impact on diagnostic assessment of therapy delivery using the Kao criteria. Further clinical studies are required to fully test the results of this initial feasibility study; however, the case of such a method for

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### Table 2

| Volume (mm³) | Static imaging | Amplitude gating |
|-------------|---------------|-----------------|
|             | Mean ± SD     | Minimum/maximum | Mean ± SD     | Minimum/maximum |
| 30 Maximum  | 2.35 ± 5.19   | 2.29/17.10      | 1.50 ± 2.63   | 0.21/89.40      |
| Region-based| 0.624 ± 0.19  | 0.20/0.84       | 0.47 ± 0.17   | 0.16/0.76       |
| Individual  | 0.12 ± 0.08   | 0.01/0.25       | 0.11 ± 0.08   | 0.008/0.23      |

Mean, SD, minimum, and maximum values are presented for each segmentation method drawn on both static and motion-corrected (amplitude gated) images.

### Table 3

| Mean dose (Gy)  | Static imaging | Amplitude gating |
|-----------------|---------------|-----------------|
|                 | Mean ± SD     | Minimum/maximum | Mean ± SD     | Minimum/maximum |
| 30 Maximum      | 82.1 ± 80.5   | 5.4/289.6       | 86.8 ± 86.9   | 8.7/313.5       |
| Region-based    | 105.0 ± 88.4  | 43.5/342.0      | 125.0 ± 102.0 | 52.6/404.0      |
| Individual lesion| 149.8 ± 101.3| 51.4/408.0      | 156.2 ± 107.1| 49.6/430.0      |

Mean, SD, minimum, and maximum values are presented for each segmentation method drawn on both static and motion-corrected (amplitude gated) images.
routine clinical use warrants publication of these initial findings.

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Some results of this work have been presented at the Society of Nuclear Medicine and Molecular Imaging over the last 2 years.

Conflicts of interest
There are no conflicts of interest.

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