Comparison of the Biograph Vision and Biograph mCT for quantitative $^{90}\text{Y}$ PET/CT imaging for radioembolisation

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Abstract

Background: New digital PET scanners with improved time of flight timing and extended axial field of view such as the Siemens Biograph Vision have come on the market and are expected to replace current generation photomultiplier tube (PMT)-based systems such as the Siemens Biograph mCT. These replacements warrant a direct comparison between the systems, so that a smooth transition in clinical practice and research is guaranteed, especially when quantitative values are used for dosimetry-based treatment guidance. The new generation digital PET scanners offer increased sensitivity. This could particularly benefit $^{90}\text{Y}$ imaging, which tends to be very noisy owing to the small positron branching ratio and high random fraction of $^{90}\text{Y}$. This study aims to determine the ideal reconstruction settings for the digital Vision for quantitative $^{90}\text{Y}$ imaging and to evaluate the image quality and quantification of the digital Vision in comparison with its predecessor, the PMT-based mCT, for $^{90}\text{Y}$ imaging in radioembolisation procedures.

Methods: The NEMA image quality phantom was scanned to determine the ideal reconstruction settings for the Vision. In addition, an anthropomorphic phantom was scanned with both the Vision and the mCT, mimicking a radioembolisation patient with lung, liver, tumour, and extrahepatic deposition inserts. Image quantification of the anthropomorphic phantom was assessed by the lung shunt fraction, the tumour to non-tumour ratio, the parenchymal dose, and the contrast to noise ratio of extrahepatic depositions.

Results: For the Vision, a reconstruction with 3 iterations, 5 subsets, and no post-reconstruction filter is recommended for quantitative $^{90}\text{Y}$ imaging, based on the convergence of the recovery coefficient. Comparing both systems showed that the noise level of the Vision is significantly lower than that of the mCT (background variability of 14% for the Vision and 25% for the mCT at $2.5\times10^3 \text{ MBq}$ for the 37 mm sphere size). For quantitative $^{90}\text{Y}$ measures, such as needed in radioembolisation, both systems perform similarly.

Conclusions: We recommend to reconstruct $^{90}\text{Y}$ images acquired on the Vision with 3 iterations, 5 subsets, and no post-reconstruction filter for quantitative imaging. The Vision provides a reduced noise level, but similar quantitative accuracy as compared with its predecessor the mCT.

Keywords: Yttrium-90, PET/CT, Radioembolisation

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Background

New digital positron emission tomography/computed tomography (PET/CT) scanners, such as the GE Discovery MI and the Siemens Biograph Vision, offer better time of flight (TOF) performance and a larger axial field of view (FOV) for higher effective sensitivity than their photomultiplier tube (PMT)-based counterparts. This increased sensitivity can be used to shorten acquisition time, reduce radionuclide activity, and/or improve image quality. It is likely that hospitals will gradually switch to these next generation digital PET/CT scanners.

The higher effective sensitivity of new digital PET/CT scanners could particularly benefit yttrium-90 ($^{90}\text{Y}$) imaging, which tends to be very noisy owing to the small positron branching ratio. PET is often regarded as the preferred imaging modality for $^{90}\text{Y}$ imaging since it offers higher resolution than Bremsstrahlung single photon emission computed tomography (SPECT) [1, 2]. It is used to image the distribution of $^{90}\text{Y}$ microspheres after radioembolisation treatment. Clinical relevant features of a post-treatment $^{90}\text{Y}$ PET scan include the lung shunt fraction (LSF), presence of extrahepatic depositions, and intrahepatic dose distribution [2, 3]. At the time of imaging, $^{90}\text{Y}$ activities in the liver typically range from 500 to 5000 MBq [4–7]. Since radioembolisation and the dosimetry involved constitute a relatively new clinical area, $^{90}\text{Y}$ imaging is often performed in the setting of research studies, where $^{90}\text{Y}$ PET is used in dose-response studies [5, 8–10]. Consistency of quantitative measurements is key throughout an imaging study, and a change of scanner during the study may affect this consistency. To warrant study consistency, a good understanding of the $^{90}\text{Y}$ imaging properties of both systems is required.

The higher sensitivity of new digital PET/CT scanners may open doors to new applications. Recently, we proposed the use of a low dosage of $^{90}\text{Y}$ microspheres (~100 MBq) for a pretreatment radioembolisation procedure for therapy planning purposes [11, 12]. Currently, pretreatment radioembolisation procedures are performed with technetium-99m ($^{99}\text{Tc}$)-labelled macro aggregated albumin (MAA). Because of the differences in shape and size between the MAA particles and the microspheres, differences between estimated dose distribution by MAA and the true dose distribution have been reported [4, 13–15]. These differences could be minimized by using the same particle for treatment and pretreatment procedure, as has been shown for holmium-166 ($^{166}\text{Ho}$) microsphere radioembolisation [16]. A safe dosage for the pretreatment $^{166}\text{Ho}$ microsphere procedure is 250 MBq [17, 18]. Inasmuch as the total energy absorbed per becquerel is higher for $^{90}\text{Y}$ than for $^{166}\text{Ho}$, this dosage would translate to 100 MBq of $^{90}\text{Y}$ to avoid unintended radiation damage [11, 12, 19].

The low activity of $^{90}\text{Y}$ for the pretreatment procedure makes imaging challenging. Using 100 MBq, Bremsstrahlung SPECT can produce quantitative images to accurately estimate the LSF, but images are of low resolution [11]. PET overestimates the LSF at 100 MBq [11]. This is caused by the low count statistics and the high random fraction, which result in a positive bias and high noise levels in the PET reconstruction. This makes PET unsuitable as an imaging modality for imaging the dose distribution of a pretreatment $^{90}\text{Y}$ procedure.

The previous study that showed the infeasibility of PET as an imaging modality for a $^{90}\text{Y}$ pretreatment procedure was performed with the PMT-based Biograph mCT PET system (Siemens) [11]. Its successor, the digital Biograph Vision, has improved spatial
resolution, improved timing resolution, an extended axial FOV, and increased sensitivity by 70% measured at the centre of the transaxial FOV (Table 1) [20]. When comparing fluorine-18 (18F) fluorodeoxyglucose images of oncological patients from both systems, the Vision scored higher in terms of overall image quality and image noise, than the mCT [22].

The improvements of the Vision with regard to the mCT are expected to lead to improved ⁹⁰Y imaging. The QUEST phantom study [23] has extensively studied the performance of multiple PET systems for ⁹⁰Y imaging and recommends reconstruction settings for quantitative purposes for the systems involved. However, the Vision was not part of the QUEST study and to our knowledge there are no recommended reconstruction settings for this system regarding quantitative ⁹⁰Y imaging.

The purpose of the present study is to evaluate the performance of the Siemens Biograph Vision in comparison with its PMT-based counterpart, the mCT, for ⁹⁰Y imaging. We used the NEMA phantom to determine the ideal reconstruction settings for the Vision in analogy to the QUEST study [23]. These reconstruction settings for the Vision were compared with the recommended reconstruction setting of the mCT [23] in terms of standardized image quality metrics. In addition, clinical relevant features for radioembolisation were compared by using an anthropomorphic phantom.

Methods

Phantoms

Two phantoms were used for this study: the NEMA image quality phantom (PTW, Freiburg, Germany) and a modified anthropomorphic thorax phantom (model ECT/TOR/P, IEL, Chilcompton, UK). The NEMA phantom is used to compare standardized image quality metrics in analogy with the QUEST study [23]. The thorax phantom is used to simulate a radioembolisation patient and to evaluate the accuracy of lung shunt estimation, extrahepatic deposition visibility, and intrahepatic activity distribution.

The NEMA phantom consists of a 9.7-L torso-shaped compartment containing six fillable spheres (inner diameter of 10, 13, 17, 22, 28, and 37 mm) and a cold, cylindrical lung insert. It was prepared in a similar way as in the QUEST study protocol [23],

| Table 1 Technical specifications of Biograph Vision and Biograph mCT |
|---------------------------------------------------------------|
| **Vision** [20] | **mCT** [21] |
| Crystals | LSO, 3.2 x 3.2 x 20 mm | LSO, 4.0 x 4.0 x 20 mm |
| Detector elements | Silicon photomultipliers | Photomultiplier tubes |
| Axial FOV | 26.1 cm | 22.1 cm |
| TOF timing resolution | 210–215 ps | 540 ps |
| Transverse spatial resolution (measured at 1 cm vertically from the centre of the FOV with ¹⁸F) | 3.7 mm (FWHM) | 4.4 mm (FWHM) |
| Sensitivity (according to NEMA NU-2 2012) | 16.4 kcps/MBq | 10.0 kcps/MBq |
| Time coincidence window | 4.7 ns | 4.1 ns |
| Energy window | 435–585 keV | 435–650 keV |
| Bed overlap | 49% | 43% |

*LSO lutetium oxyorthosilicate, FOV field of view, FWHM full width at half maximum*
where the phantom was filled with $^{90}$Y chloride, in 0.5 M HCl to prevent adhesion to the plastic phantom walls [24], to acquire a sphere-to-background concentration ratio of 8 and a total initial activity of $2.5 \times 10^3$ MBq.

The modified thorax phantom consists of a torso-shaped compartment containing a liver compartment with a solid tumour (sphere of 15.9 mL) and a necrotic tumour (outer rim 18.9 mL and inner sphere 5.6 mL), two lung compartments, a cylindrical spine insert, and three extrahepatic compartments (spheres of 2.0, 4.2 and 8.2 mL). The phantom was filled with $^{90}$Y chloride in 0.5 M HCl, to acquire a lung shunt fraction (LSF) of 5.0%, a tumour to non-tumour ratio (T/N) of 8.0, and a total initial activity of $1.0 \times 10^3$ MBq. The activity concentration of the extrahepatic depositions in the phantom was based on the median size (6.8 mL, range 1.1–42 mL) and median activity (1.3% of the infused activity) of 34 extrahepatic depositions found by Prince et al. [18] and was therefore chosen to be 1.3% of the total activity in the phantom/6.8 mL.

**Image acquisition**

Both phantoms were scanned consecutively on a Siemens Biograph mCT and a Siemens Biograph Vision. Table 1 lists technical specifications of both systems. Scans were acquired during decay and all activities at the time of imaging are listed in Table 2.

Both phantoms were scanned in two bed positions of 15 min per bed position on both systems. A CT scan was made for attenuation correction and to support delineation.

**Image reconstruction**

Images acquired by the mCT were reconstructed with the reconstruction settings recommended by the QUEST study [23]. This is an ordered subset expectation maximization (OSEM) reconstruction algorithm, including time of flight (TOF) information, resolution recovery (TrueX), attenuation, scatter, random, dead time, and decay correction. The reconstruction used 2 iterations with 21 subsets, resulting in a voxel size of $4.1 \times 4.1 \times 3.0$ mm$^3$, and no post-reconstruction filter was applied.

Since, to our knowledge, no prior study has been published on $^{90}$Y PET imaging using the Vision, images acquired at the Vision were reconstructed with a variety of reconstruction settings, so as to determine the optimal setting. All reconstruction methods used an OSEM reconstruction algorithm, including TOF information, resolution recovery (TrueX), attenuation, scatter, random, dead time, and decay correction. The number of subsets is fixed at 5 by the vendor and the number of iterations was varied between 1 and 17. Images were reconstructed on a $220 \times 220$ matrix resulting in a voxel size of $3.3 \times 3.3 \times 3.0$ mm$^3$. No post-reconstruction filter was applied. The

| Table 2 | Total activity (MBq) in the NEMA and thorax phantoms at time of imaging |
|---------|-------------------|---|---|---|---|---|---|---|
|         | Day 0 | Day 2 | Day 4 | Day 6 | Day 8 | Day 9 | Day 10 |
| NEMA mCT | 2511  | 1560  | 901   | 537   | 328   | 259   | 194    |
| Vision  | 2563  | 1526  | 930   | 553   | 322   | 264   | 205    |
| Thorax mCT | 961  | 597   | 345   | 206   | 126   | 99    | 74     |
| Vision  | 982   | 585   | 356   | 212   | 123   | 101   | 78     |
optimal reconstruction was determined based on convergence of the recovery coefficient with iteration number.

**Analysis**

For the NEMA image quality phantom, we analysed the same metrics as the QUEST phantom study. These are:

Percent background variability (BV), following the NEMA NU 2-2007 guidelines, defined for each sphere size as:

\[
BV = \frac{SD_{B,s}}{C_{B,s}} \times 100\%
\]

(1)

where \(SD_{B,s}\) is the standard deviation of the average of the 60 background regions of interest (ROIs) for sphere size \(s\) and \(C_{B,s}\) is the average count of the 60 background ROIs for sphere size \(s\).

Percentage misplaced counts in the lung insert (\(\Delta C_{\text{lung}}\)), following the NEMA NU 2-2007 guidelines, defined as:

\[
\Delta C_{\text{lung}} = \frac{C_{\text{lung}}}{C_B} \times 100\%
\]

(2)

where \(C_{\text{lung}}\) is the average count in the lung insert ROI and \(C_B\) is the average count of the background ROIs.

Background concentration accuracy (BCA), defined for each sphere size as:

\[
BCA = \frac{a_{B,\text{measured}} - a_{B,\text{true}}}{a_{B,\text{true}}} \times 100\%
\]

(3)

where \(a_{B,\text{measured}}\) is the measured activity concentration in the background and \(a_{B,\text{true}}\) is the true activity concentration in the background.

Total activity accuracy (TAA), defined as:

\[
TAA = \frac{A_{\text{TOT,measured}} - A_{\text{TOT,true}}}{A_{\text{TOT,true}}} \times 100\%
\]

(4)

where \(A_{\text{TOT,measured}}\) is the measured total activity in the entire FOV and \(A_{\text{TOT,true}}\) is the known true total activity in the phantom.

Recovery coefficient (RC), following the NEMA NU 2-2007 guidelines for delineation of the spheres, defined for each sphere size as:

\[
RC = \frac{a_{S,\text{measured}}}{a_{S,\text{true}}} \times 100\%
\]

(5)

where \(a_{S,\text{measured}}\) is the measured activity concentration in the sphere and \(a_{S,\text{true}}\) is the true activity concentration in the sphere.

To evaluate the thorax phantom, we assessed the following metrics, which are commonly assessed for radioembolisation treatment planning:

The lung shunt fraction (LSF), defined as:

\[
LSF = \frac{C_{\text{lung}}}{C_{\text{lung}} + C_{\text{liver}}} \times 100\%
\]

(6)

where \(C_{\text{lung}}\) is the total count in the lungs dilated by 6 mm and \(C_{\text{liver}}\) is the total count in the liver dilated by 6 mm. The lung and liver contours were dilated by the
spatial resolution of the PET systems to partially compensate for the partial volume effect.

Tumour to non-tumour ratio (T/N) of the solid and the necrotic tumour, defined as:

\[
\frac{T}{N} = \frac{C_{\text{tumour}}}{C_{\text{parenchyma}}} 
\]  

where \( C_{\text{tumour}} \) is the average count in the tumour volume of interest (VOI) and \( C_{\text{parenchyma}} \) is the average count in the parenchymal VOI.

Parenchymal dose per GBq of \(^{90}\)Y administered, defined as:

\[
D_{\text{parenchyma}} = \frac{A_{\text{parenchyma}}}{A_{\text{total}}} \times 50 
\]

where \( A_{\text{parenchyma}} \) is the activity in the parenchymal VOI, \( A_{\text{total}} \) is the known total activity in the phantom, \( m_{\text{parenchyma}} \) is the mass of the parenchymal VOI (determined using the liver VOI volume and a conversion factor of 1.03 g/mL), and 50 is the absorbed energy in joule from the decay of 1 GBq of \(^{90}\)Y.

Contrast-to-noise ratio (CNR) of the extrahepatic depositions as a measure of visibility, defined as:

\[
\text{CNR} = \frac{C_{\text{deposition}} - C_{\text{B}}}{\text{SD}_B} 
\]

where \( C_{\text{deposition}} \) is the average count in the extrahepatic deposition VOI, \( C_{\text{B}} \) is the average count in a background VOI, and \( \text{SD}_B \) is the standard deviation of a background VOI.

Differences between the Vision and the mCT are tested for significance with a paired t-test assuming a 5% significance level.

Results

Figure 1 shows the total prompts, randoms, and net trues for the Vision and mCT acquisitions of the NEMA and thorax phantoms. The net trues show a strong linear trend with activity, of which the slope, intercept, and coefficient of determination are listed in Table 3. The Vision has an increased trues rate compared with the mCT by 68% (NEMA phantom) and 65% (thorax phantom).

Ideal reconstruction setting Vision

Figure 2 shows the transverse slice of the NEMA phantom at day 0 (2.5·10^3 MBq) for three reconstruction settings for the Vision acquisition and for the 2-iterations setting for the mCT acquisition. Visually, the reconstructions of the Vision outperform the reconstruction of the mCT because of the lower noise level and the better visibility of the (smaller) spheres. Increasing the number of iterations increases noise.

Figure 3 shows the recovery coefficient and the percentage background variability plotted against iteration number of the different sphere diameters at day 0. Depending on the study purpose, one can either choose a number of iterations that favours high recovery (higher number of iterations) or one that favours low noise (low number of iterations). Since the purpose of this study is quantitative imaging, we like to achieve a
The RC curves of the three largest spheres shown in Fig. 3 converge at 3 iterations. The RC curves of the three smallest spheres show more varying trends. In order to not increase the background variability while the recovery coefficient barely improves, we choose 3 iterations with 5 subsets as the ideal reconstruction setting for the Vision. For subsequent comparisons between the Vision and the mCT, the images acquired by the Vision will therefore be reconstructed using 3 iterations with 5 subsets without a post-reconstruction filter.

**Comparison Vision and mCT**

Figure 4 compares the NEMA phantom scanned at the Vision (reconstructed with 3 iterations, 5 subsets) with the mCT scans (reconstructed with 2 iterations, 21 subsets). Paired *t* tests showed a significant difference between the Vision and the mCT for background variability (BV) (*p* value < 0.001) and background concentration accuracy (*p* value < 0.05). The BV was about twice as high for the mCT. This affects the visibility of hot and cold spots. For other quantitative measures, there was no significant difference between the Vision and the mCT.

The thorax phantom mimics a radioembolisation patient and has an activity distribution that is clinically more relevant than that of the NEMA phantom. Figure 5 shows

| Measurement       | Slope (counts/MBq) | Intercept (counts)        | $R^2$  |
|-------------------|--------------------|--------------------------|--------|
| NEMA Vision       | 448 [439–457]      | 1.57·$10^4$ [5.16·$10^3$–2.62·$10^4$] | 0.9997 |
| NEMA mCT          | 266 [261–271]      | 1.04·$10^4$ [4.53·$10^3$–1.62·$10^4$] | 0.9997 |
| Thorax Vision     | 263 [259–267]      | 1.58·$10^4$ [1.37·$10^3$–1.78·$10^4$] | 0.9998 |
| Thorax mCT        | 159 [152–167]      | 0.762·$10^4$ [4.15·$10^3$–1.11·$10^4$] | 0.9983 |
metrics that are commonly assessed for radioembolisation treatment planning. Paired t tests showed a significant difference between both systems for the LSF (p value < 0.05), the parenchymal dose (p value < 0.05), and the CNR (p value < 0.001). Although significant differences were observed, both systems showed similar trends for these metrics. Both overestimated the LSF, and as expected, the overestimation got worse when the total activity decreased. This can be attributed to the earlier described positive bias in maximum likelihood reconstructions at low activities [11, 25]. However, unexpectedly, the Vision showed a slightly larger overestimation of the LSF than the mCT. Metrics regarding intrahepatic dose distribution (T/N and parenchymal dose) were underestimated by both systems and showed large variations. The CNR, a measure for the visibility of the extrahepatic depositions, showed the most stable trend and decreased with decreasing phantom activity. The CNR was considerably higher for the Vision than for the mCT.

**Discussion**

In this study, we evaluated the performance of the Vision for ⁹⁰Y imaging and compared it with the performance of its predecessor, the mCT. We determined the optimal reconstruction settings for the Vision to be 3 iterations with 5 subsets. We found that the Vision outperformed the mCT in terms of noise level, but performed similarly in terms of quantitative accuracy.
The prompts, randoms, and trues of both the Vision and the mCT show good linearity, which implies that no saturation of the detectors occurs. The Vision had 65% (thorax phantom) or 68% (NEMA phantom) more net trues than the mCT. This is in agreement with the 70% increase in sensitivity found by van Sluis et al. for $^{18}$F [20].

The slope of the true count with activity is dependent on the geometry and activity distribution of the scanned object, since the slopes of the true count with activity on the same scanner with the same acquisition parameters differ between NEMA and thorax phantoms. The total prompts and randoms have an offset due to the presence of lutetium-176 ($^{176}$Lu) in the crystals. The magnitude of this offset is dependent on, e.g., the amount of crystals (and therefore the amount of $^{176}$Lu), the number of possible coincidence line-of-response, the time coincidence window, and the energy window [26]. All these specifications differ between the mCT and the Vision and result in a larger offset for the mCT than for the Vision.

![Fig. 4 Results of the NEMA phantom comparing the mCT (2 iterations, 21 subsets) with the Vision (3 iterations, 5 subsets). BCA, background concentration accuracy](image-url)
Scans at both systems were acquired in step-and-shoot mode, instead of continuous-table-motion mode. For scans with a short scan length and long acquisition time, like ⁹⁰Y PET liver scans, it has been shown that step-and-shoot mode has a higher counting efficiency compared to continuous-table-motion mode [27]. Therefore, we chose to perform all scans in step-and-shoot mode to obtain the best image quality and quantification precision.

The ideal reconstruction setting is dependent on the study purpose. Just like the QUEST study, we focused on quantitative accuracy. We recommend to reconstruct ⁹⁰Y data acquired on the Vision with 3 iterations, 5 subsets, and no post-reconstruction filter. This is based on the fact that after three iterations quantitative metrics will only marginally improve while noise will still substantially increase. When the objective of a scan is focused on visual image quality, one could decide to add a post-reconstruction filter. In this study, we did not study the influence of different filters in detail, since Gaussian filters will reduce quantitative accuracy.

The results from our NEMA measurements on the mCT were comparable with the results from the QUEST study. There were some small deviations (like the steep trend
seen for our background variability versus sphere diameter, which is less pronounced in the QUEST study, but these can be explained by the fact that the data represented by QUEST consisted of multiple scanners. The measurements on our single scanner still fall within the error margins of the QUEST study.

Comparing quantitative results of the Vision with the mCT does not show clinically relevant differences between the systems. This suggests that a smooth upgrade during quantitative studies is possible, if the proper reconstruction protocol is chosen. However, the noise level of the Vision is substantially lower than the noise level of the mCT. This could make visualization of hot and/or cold regions, such as extrahepatic depositions, easier on the Vision. van Sluis et al. shows similar results comparing the Vision and the mCT for $^{18}$F, where they show that the Vision outperforms the mCT visually, but based on quantitative measures, both systems are comparable [22].

For the specific clinical case of $^{90}$Y imaging for radioembolisation, quantitative measures are important. Although the LSF and parenchymal dose show significant differences between the Vision and the mCT, these differences are small (mean LSF difference is 0.9% point (pp), mean parenchymal dose difference is 2.8 Gy/GBq). Furthermore, the uncertainty in volume measurement and uncertainty in the activity calibration cause an uncertainty of ±10% in the true parenchymal dose of the thorax phantom. This makes it difficult to conclude with certainty whether the Vision or the mCT results in a more accurate estimated parenchymal dose. With a $^{90}$Y pretreatment procedure in mind, errors in estimated parenchymal dose could cause under- or overdosage. In combination with the overestimation of the LSF by 9 pp at 100 MBq, we have to conclude that the Vision is, just like its predecessor the mCT, unsuitable as an imaging system for imaging of a theoretically safe pretreatment dosage of 100 MBq $^{90}$Y.

The Vision does have a lower noise level compared with the mCT. This means that the Vision would be the more appropriate system when addressing the dose distribution visually, for example after radioembolisation treatment to identify whether the dose distribution in a tumour is homogenous or not.

**Conclusion**

In this study, we evaluated the performance of the Vision for $^{90}$Y imaging and compared it with the performance of its predecessor, the mCT. We recommend to reconstruct $^{90}$Y images acquired on the Vision with 3 iterations, 5 subsets, and no post-reconstruction filter, for quantitative purposes. Visually, the Vision outperforms the mCT because of its lower noise level, but based on quantitative measurements, both scanners perform similarly.

**Abbreviations**

ΔC$_\text{lung}$: Percentage misplaced counts in the lung insert; $^{166}$Ho: Holmium-166; $^{176}$Lu: Lutetium-176; $^{18}$F: Fluorine-18; $^{90}$Y: Yttrium-90; $^{99m}$Tc: Technetium-99m; BCA: Background concentration accuracy; BV: Background variability; CNR: Contrast-to-noise ratio; FOV: Field of view; FWHM: Full width at half maximum; LSF: Lung shunt fraction; LSO: Lutetium oxyorthosilicate; MAA: Macro aggregated albumin; OSEM: Ordered subset expectation maximization; PET/CT: Positron emission tomography/computed tomography; PMT: Photomultiplier tube; pp: Percentage point; RC: Recovery coefficient; ROI: Region of interest; SPECT: Single-photon emission computed tomography; T/N: Tumour to non-tumour ratio; TAA: Total activity accuracy; TOF: Time of flight; VOI: Volume of interest

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Authors’ contributions
BK acquired the scans, analysed the data, and drafted the manuscript. CB contributed to the design of the study and the interpretation of results. ML and MV contributed to the design of the study. HdJ contributed to the design of the study and the interpretation of results. All authors read and approved the final manuscript.

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Availability of data and materials
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Consent for publication
Not applicable.

Competing interests
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