First Phantom-Based Quantitative Assessment of Scandium-44 Using a Commercial PET Device

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Scandium (Sc) is a promising candidate for theranostic applications due to the existence of radioisotopes suitable for both imaging and therapy. A “proof-of-concept” study regarding first-in-human use of $^{44}$Sc for imaging metastatic neuroendocrine tumors was reported recently, however, quantitative assessment of $^{44}$Sc-based PET images was not performed. The aim of this study was to evaluate quantitative capabilities of $^{44}$Sc-PET using a commercial PET scanner. The NEMA/IEC body phantom with $^{44}$Sc was acquired according to the local protocol used for whole-body oncological $^{18}$F-FDG PET examinations. Additionally, we characterized the signal recovery (recovery coefficient—RC) according to the iteration number. For all reconstructions, pertinent image corrections (normalization, dead time, activity decay, random coincidence, and attenuation) were applied. Presently, $^{44}$Sc scatter corrections are not optimized and could, thus, result in quantitative bias. To investigate the best option, the data were reconstructed using different available scatter corrections (relative -RelSC- and absolute -AbsSC-) and an additional prompt-gamma correction (PGC). System cross-calibration with the local dose calibrator ($\text{BG}_{\text{cal}}$) and image noise, expressed by the coefficient of variation (COV), were evaluated in the homogeneous background region (5 kBq/mL) of the phantom. Maximum ($\text{RC}_{\text{max}}$) and 50% threshold recovery coefficients, corrected for background ($\text{RC}_{\text{A50}}$), were measured for all spherical inserts (25 kBq/mL) of the phantom. Acceptable COV (<15%) was achievable with low iteration numbers (<3). $\text{BG}_{\text{cal}}$ differences were low: mean $\text{BG}_{\text{cal}}$ were 77.8, 81.3, and 86.7%, for RelSC, AbsSC, and PGC, respectively. RC values exceeded the present RC range recommended for $^{18}$F-FDG procedures. Using the iterations to be evaluated, $\text{RC}_{\text{A50}}$ ranged from 29.9 to 59.9% for the smallest lesion (spherical insert of 10 mm diameter) and from 45.5 to 80.3% (13 mm), 66.4 to 75.6% (17 mm), 71.7 to 75.7% (22 mm), 75.1 to 78.6% (28 mm), and 76.7 to 80.9% (37 mm) for the, respectively spherical inserts. The results of
1. INTRODUCTION

Clinical theranostic application of $^{68}$Ga ($T_{1/2} = 68$ min, $E_\beta^{+} = 830$ keV, $I = 89\%$) and $^{177}$Lu ($T_{1/2} = 6.65$ d, $E_\beta^{-}$ av = $134$ keV, $I = 100\%$; $E_\gamma$ = $113$ keV $I = 6\%$, $208$ keV $I = 10\%$) is now common for PET imaging and targeted radionuclide therapy, respectively. The $^{68}$Ga/$^{177}$Lu theranostic couple is used for labeling with somatostatin analogs (e.g., DOTATOC or DOTATATE) and, more recently, also for PSMA-targeting ligands (e.g., PSMA-11 and PSMA-617), thereby, enabling radiotheranostics of neuroendocrine and prostate cancer, respectively [1–4].

PET is currently regarded as more sensitive technique than SPECT and provides images with superior spatial resolution and improved quantification [5]. The switch from $^{111}$In-based SPECT imaging, such as $^{111}$In-octreotide, to the use of $^{68}$Ga for PET imaging, represents a cornerstone in nuclear medicine: $^{68}$GaGa-DOTATATE demonstrated significantly superior image quality and, hence, outperformed $^{111}$In-octreotide in both diagnostic accuracy and impact on treatment decisions [6].

$^{68}$Ga is obtained from a $^{68}$Ge/$^{68}$Ga-generator system, which can be conveniently placed in a radiopharmacy at any hospital. The availability of $^{68}$Ga is, however, restricted with regard to activity that can be eluted for radiopharmaceutical preparation, while the generator itself is also costly. A drawback of using $^{68}$Ga, however, is its short half-life of only $68$ min (compared to $2.81$ d for $^{111}$In), which does not allow scanning at late time points after injection nor transportation of $^{68}$Ga-based radiopharmaceuticals over longer distances (when a $^{68}$Ge/$^{68}$Ga generator is not present on site). Optimally, the physical half-life is identical to the biological half-life and $^{44}$Sc can be used with targeting agents that are retained longer in the body. As an example, $^{44}$Sc may be viable in the use of albumin-binding radiopharmaceuticals, which have longer circulation time [7]. Finally, it remains to be mentioned that $^{68}$Ga has a different coordination chemistry compared to $^{177}$Lu, therefore, $^{68}$Ga results in different chelator-radiometal complexes and, hence, chemically unequal radiopharmaceuticals. This may result in different in vivo uptake patterns for both diagnostic and therapeutic radioligands [8–10].

Several years ago, $^{44}$Sc was proposed as an alternative PET nuclide [11, 12]. The physical decay properties [13] of $^{44}$Sc ($T_{1/2} = 4.04$ h, average $E_\beta^{+} = 632$ keV, $I = 94\%$) are attractive to address the shortcomings of $^{68}$Ga mentioned above. It decays with an almost four-fold longer half-life than $^{68}$Ga [13] by emitting relatively low energy $\beta^{+}$-particles that enables the acquisition of PET images with a spatial resolution equal or even superior to that achievable with $^{68}$Ga [14]. The coordination chemistry allows the preparation of $^{44}$Sc-radioconjugates using a DOTA-chelator in analogy with $^{177}$Lu-labeling [9]. Preclinical experiments in mice showed almost identical distribution patterns for $^{44}$Sc- and $^{177}$Lu-labeled ligands [10, 11]. Consequently, $^{44}$Sc may be adopted as a more favorable diagnostic match to $^{177}$Lu than $^{68}$Ga. In future, $^{47}$Sc ($T_{1/2} = 3.35$ d, average $E_\beta^{-} = 162$ keV, $I = 100\%$; $E_\gamma$ = $159$ keV $I = 68\%$) can be employed as a therapeutic match to provide chemically-identical $^{44}$Sc/$^{47}$Sc-based radiopharmaceuticals for radiotheranostic applications [15–17]. A possible addition to the scandium radioisotope family, due to its theoretically more favorable characteristics for PET imaging, is $^{45}$Sc [18]. Although it provides possible advantages in terms of gamma emission, it is currently more challenging to produce in terms of yield, as well as cost [19, 20].

Rösch and co-workers performed preliminary studies using $^{44}$Sc obtained from a $^{44}$Ti/$^{44}$Sc generator system [21]. First PET images of a patient administered with $^{[44]$Sc]$^{[44]$Sc-DOTATOC looked promising, where somatostatin-receptor-positiver liver metastases were imaged and visible $18$ h after administration of $37$ MBq [22]. It was later proposed to produce $^{44}$Sc at a cyclotron using natural Ca targets, in an attempt to simplify production and increase yield [23, 24]. At PSI, it was demonstrated for the first time that $^{44}$Sc can also be produced in high quantities and good quality when using enriched $^{44}$Ca targets [11, 25]. In recent years, the production was constantly improved, currently enabling the preparation of up to $5$ GBq $^{44}$Sc with small quantities of target material. This production method may be translated to a conventional medical cyclotron commonly installed in PET centers. Subsequently, the application of cyclotron-produced $^{44}$Sc in two patients with neuroendocrine neoplasms, receiving a diagnostic administration of $^{[44]$Sc]$^{[44]$Sc-DOTATOC at Zentralklinik Bad Berka (Germany), was reported [26]. First-in-man studies with $^{[44]$Sc]$^{[44]$Sc-PSMA-617, using generator-produced $^{44}$Sc, were reported recently in the literature, underlining the excellent features of $^{44}$Sc in a clinical setting [27].

Despite the reported clinical advantages of $^{44}$Sc, its use could potentially be restricted by the currently limited quantification capability of clinical devices. The first reported human study using $^{44}$Sc [26] indicated that absolute quantification was not available, but the authors did not discuss this limitation further. Other studies on the use of non-pure positron emitters started evaluating this limitation. In particular, the importance of adopting adapted and optimized corrections in the acquisition and reconstruction process, in achieving quantitative imaging in clinical setups, was highlighted. As an example, the work by Armstrong et al. [28] addressed the impact of prompt-gamma coincidence inducing quantitative artifacts and underlined the important role of specific corrections in absolute quantification using $^{82}$Rb cardiac imaging. The study by Lubberink and Herzog

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[29] reviewed different optimization strategies for improving quantification of $^{124}$I and $^{86}$Y, including effects from scatter and prompt-gamma corrections.

The aim of this paper was to evaluate the quantification capabilities of $^{44}$Sc in a clinically-relevant phantom configuration study performed with a commercial PET scanner.

### 2. MATERIALS AND METHODS

The experimental design presented in this work included (a) image acquisitions of a NEMA/IEC NU2 phantom (see sections 2.1, 2.2) and (b) the subsequent data analysis (see sections 2.3, 2.4). The phantom was scanned under clinically-relevant conditions at the Aarau Cantonal Hospital (KSA) using a mCT Biograph PET-CT (Siemens Healthineers AG) [30]. $^{44}$Sc was produced via the $^{44}$Ca(p,n)$^{44}$Sc nuclear reaction at Injector 2 cyclotron facility at Paul Scherrer Institute (PSI), Switzerland, as previously reported [25, 26].

The reference activity received from PSI was used to perform a cross calibration of the local dose calibrator (Veenstra Instruments, Joure, Netherlands VDC-405 with ionization chamber VIK-202). At PSI, the nuclide content in the irradiated targets was identified and quantified by $\gamma$-ray spectrometry using an N-type high-purity germanium (HPGe) coaxial detector (EURISYS MESURES, France) and the Ortec InterWinner 5.0 software. A dose calibrator (ISOMED 2010, Nuclear-Medizintechnik Dresden GmbH, Germany) was employed for the quantitative determination of $^{44}$Sc activity [20]. The reference activity was measured at the end of chemical separation and the time noted, before packing and transportation. This reference activity and time, on its original vial, was used to cross calibrate the local dose calibrator by modifying the isotope settings (factor and scale) to match this reference activity. The factor obtained was then used for all dispensing syringe activities. The activities obtained for various settings, along with the setting chosen for $^{44}$Sc, are listed in Table 1. All measurements were decay-corrected to the reference activity. These values were then used to estimate the injected activity (IA) used to fill the main phantom volume (background volume) and the included spherical inserts, representing lesions. The $^{44}$Sc cross-calibrated factor (presented in Table 1) was then used for all activity dispensing.

### Table 1 | Reference activity and employed factors for determination of the local dose calibrator factor for $^{44}$Sc measurements.

| Radionuclide | Setting (Factor and scale) | Measurement (GBq) |
|--------------|----------------------------|-------------------|
| Ga-68        | 751 scale 1                | 2.116             |
| F-18         | 765 scale 1                | 1.401             |
| Co-60        | 889 scale 1                | 0.392             |
| Sc-44        | 760 scale 0.56             | 0.594             |
| Reference activity |                      | 0.595             |

All measurements were decay-corrected to the reference time provided by PSI. The factor is constrained to a given range (0–900) and for some radionuclides, a scale different than 1 is also needed.

2.1. Phantom Preparation

The NEMA/IEC NU2 phantom (PTW, Freiburg, Germany) used in this study enabled the characterization of image quality and quantitative signal recovery in the main background phantom volume and in spherical structures of different size potentially affected by partial-volume-effect (PVE). The NEMA/IEC phantom is composed of a main volume (background) of 9.5 L that mimics the human abdominal shape. The phantom includes a set of hollow-glass-sphere inserts of variable diameters: 10, 13, 17, 22, 18, and 37 mm, respectively. In addition, a lung insert (5-mm-diameter and 16-cm-long cylinder filled with plastic material to reproduce the lung density of 0.3 g/mL) was placed in the center of the phantom to simulate lung tissue attenuation. The phantom was filled with a background activity concentration of 5 kBq/mL and a five-fold higher activity concentration (25 kBq/mL) in the spherical inserts. The activity concentration and background ratio used was chosen to mimic the average hepatic activity concentration measured in patients when using $^{18}$F-FDG oncological PET 1 h after administration of a mass-activity of 3.5 MBq/kg. This corresponded to the recommended dose reference level in Switzerland at the time of this study for this specific examination [31]. The adopted experimental setup allowed comparison with previously published results obtained with $^{18}$F-FDG for the same device [31].

2.2. PET Acquisition and Reconstructions

The Siemens mCT PET, utilized for the phantom acquisitions performed in this work, consists of 4 rings, with 48 detector blocks in each ring and lutetium oxyorthosilicate (LSO) crystals, a PET field-of-view (FOV) of 22.1 cm and transaxial FOV of 70 cm. The detector ring diameter is 84.2 cm, the reported time coincidence window is 4.1 ns, system time-of-flight (TOF) 540 ps, and energy window 435–650 keV. The phantom was placed on the PET bed to have the equatorial plane of the spherical inserts at the center of the device FOV, where the system sensitivity is expected to be at its maximum. A single-bed, step-and-shoot, 300 s, list-mode (LM) acquisition was performed. The LM data were reconstructed according to the local clinic protocol used for whole-body oncological $^{18}$F-FDG PET examinations, with acquisition time of 150 s per bed position. For the acquisition, the vendor specific setup for the radionuclide $^{44}$Sc was selected, taking a positron fraction of 0.9427 and half-life of 3.97 h into consideration.

Supplementary reconstructions were performed, in addition to the local clinical protocol, by varying the number of iterations from 1 to 10 with 21 fixed subsets. This was done in order to characterize the convergence of the signal recovery in background areas, the spheres, and the lung. Reconstructions were performed using the TOF information and the point-spread function correction implemented in the vendor-based iterative reconstruction algorithm (TrueX + TOF - ultraHD-PET).

2.3. Data Analysis

2.3.1. PET vs. Dose Calibrator Activity Cross-Calibration ($BG_{cal}$)

The PET to local dose calibrator activity cross calibration ($BG_{cal}$) was tested by calculating the ratio between the measured PET
signal ($\bar{A}_{c, bg}$) and the expected average activity concentration ($A_{c, bg}$) evaluated in the homogeneous background regions of the phantom:

$$BG_{cal} = \frac{\bar{A}_{c, bg}}{\bar{A}_{c, bg}}$$

(1)

$\bar{A}_{c, bg}$ was measured in four cubic regions of interest (40 mm per side) placed in the homogeneous background region of the phantom which surrounded the spheres. When using $^{18}$F]FDG, a deviation $<0.1$ from the ideal $BG_{cal} = 1$ is regarded as acceptable.

2.3.2. Noise

The image noise was evaluated by the coefficient of variation (COV), which is the ratio between the standard deviation and the average activity concentration measured in same cubic volumes of interest (VOIs) of the phantom background described above for the cross-calibration assessment:

$$COV(\%) = \frac{SD_{bg}}{\bar{A}_{bg}} * 100$$

(2)

The background signal-to-noise ratio (SNR) is the reciprocal of the COV. A COV $\leq 15\%$ (background SNR $\geq 6.7$) was considered as an acceptable noise level for clinical image interpretation, as suggested in the EARL procedure [32]. Although this value is somewhat arbitrary, it has already been used as a reference value in previously-published works [33–35], which enables a term of comparison for image quality assessments.

2.3.3. Average Residual Lung Error

A cylindrical VOI 30 mm in diameter and 120 mm in length was included in the lung insert. The average residual lung error (in %) was obtaining by:

$$LE(\%) = \frac{\bar{L}_c}{\bar{A}_{c, bg}} * 100$$

(3)

where $\bar{L}_c$ is mean counts in the lung region and $\bar{A}_{c, bg}$ is the average activity concentration in the background region.

2.3.4. Recovery Coefficient

A VOI, with a side of 50 mm in length, was centered on each of the six spheres of the phantom. For each spherical insert ($j = 1...6$) the maximum and the background-adapted recovery coefficients (RC) were obtained as follows:

$$RC_{j,max}(\%) = \frac{a_{c,sph,j,max}}{A_{c,sph}} * 100$$

$$RC_{j,A50}(\%) = \frac{a_{c,sph,A50}}{A_{c,sph}} * 100$$

(4)

(5)

where $A_{c,sph}$ is the expected activity concentration in the spheres, $a_{c,sph,j,max}$ is the measured maximum voxel value (in terms of activity concentration) for a given spherical insert. $a_{c,sph,A50}$ is, for each considered spherical insert, the average voxel value in a VOI defined by a 3D iso-contour adapted for background [as defined in [32]] as recommended by the EANM Guidelines for $^{18}$F]FDG tumor PET imaging [35]. The numerator of Equation (5), for a given sphere $j$, is the average activity concentration computed from voxels in the VOI$_{sph,A50}$. The VOI$_{sph,A50}$ includes all voxels with activity concentration $\geq (a_{cj, max} + \bar{A}_{c, bg})/2$, where $a_{cj, max}$ is a maximum activity concentration assumed in the hot sphere $j$. RC calculations were performed using a Matlab script.

Calculated $RC_{max}$ and $RC_{A50}$ values were compared with reference values provided by the EANM/EARL accreditation protocol [36].

2.3.5. Evaluation of Available Scatter and Prompt Gamma Corrections

Pertinent image corrections (normalization, dead time, activity decay, random coincidence, and attenuation) were applied to all reconstructions performed. A known limitation that applies to radionuclides not routinely used in clinical studies (particularly if they are non-pure positron emitters) exists on the applied system corrections that were optimized for the $^{18}$F PET setting and not, in this case, for the $^{44}$Sc.

For this reason, along with the clinical corrections, all possible scatter corrections available in the iterative reconstruction algorithm (relative and absolute scatter corrections) were tested, thereby, producing a set of dedicated reconstructions [37]. Additionally, we applied the available prompt-gamma correction (PGC) [38] to evaluate the contributions of high-energy $\gamma$-emissions of $^{44}$Sc to the quantitative accuracy of reconstructed PET images.

Evaluated scatter methods differ mainly on the procedure used to account for scatter events at the sinogram level [39]. The absolute scatter correction relies on the single scatter estimation (SSS) algorithm [37], which estimates the scatter distribution produced by a single scatter event and integrates this contribution over the whole acquired FOV volume. The relative scatter correction, on the other hand, scales the SSS (plane-by-plane) to the net trues based on a scaling mask computed outside the emission object. The volume covered by the emission object is known from the transmission sinogram obtained from the CT acquisition. Based on the CT information, the emitting object and the outside region masks are generated and applied to correct the emission object sinogram. Similar to the methodology, the PGC is estimated by the single event rates. The PG model is combined with a scatter sinogram estimated by the SSS algorithm, and the total background model is scaled to fit the tails of the measured data [38]. Additionally, further contributions from PGC can be addressed as described by Armstrong et al. [28] and Lubberink and Herzog [29].

Statistical analyses were performed with Python Scipy statistical module using the Wilcoxon signed rank test. Additionally, differences between the obtained COV and BG$_{cal}$ were calculated between the reconstructions obtained in
comparison to the original/reference reconstruction (absolute scatter correction with PSF and TOF).

\[
\text{Difference}(\%) = \frac{S_{\text{refrecon}} - S_{\text{testedrecon}}}{S_{\text{refrecon}}} \times 100 \quad (6)
\]

### 3. RESULTS

The effect of the iteration number on the noise (COV), average residual lung error and the BG calibration obtained in the tested reconstructed images are presented in Figure 1.

Noise level within acceptable ranges (15% was used as level of comparison to \[^{18}\text{F}]\text{FDG} [32]) is achievable with low iteration numbers (iterations \(\leq 3\) when subsets = 21) and increases as the iteration number increases for all evaluated corrections.

Figure 1 also shows the effect of the used scatter correction (SC) model (absolute vs. relative) on the obtained COV and background calibration values. The main observed result is the reduced background calibration level obtained for reconstructions employing the relative scatter in comparison to other evaluated corrections. The BG calibration values did not vary with the tested number of iterations, averaged at 0.81 (± 0.01), 0.78 (± 0.01), and 0.87 (± 0.01), respectively for the absolute scatter correction, relative scatter correction and PGC. With the results obtained with PGC, it was observed that the BG calibration improved in relation to the original (absolute scatter correction and no PGC), and the previously-evaluated relative scatter. Both differences were statistically significant with \(p\)-value < 0.001. Figure 1 also shows the average residual lung error obtained.

Additionally, Figure 1 shows that the average residual lung error reduces with the increase of iteration number and a small difference (≤ 4%) was observed between the evaluated corrections.

Figure 2 shows scatter corrections and PGC profiles obtained on the acquired sinogram. All plots discriminate the contributions from the emission and scatter estimate sinograms and the subtraction mask used for scatter modeling without (B–relative scatter) and with PGC (C), respectively. Differences depends on the size of the applied scatter mask (blue dashed line) and, subsequently, the extrapolation of events obtained outside the central part of the sinogram and the scaled scatter estimate (green line). Noticeably, the initial scatter sinogram (SSS) is further overestimated with the relative scatter by using the wider mask, the PGC is able to improve the applied correction. The effect of this was seen in the BG calibration in Figure 1.

Lastly, recovery coefficients were evaluated for all reconstructed PET images, depending on the applied correction (PGC, no PGC with absolute scatter correction and no PGC with relative scatter correction), as a function of the number of iterations. Figure 3 shows the obtained \(R_{C_{50}}\) and \(R_{C_{\text{max}}}\) for all spherical inserts of the phantom across the evaluated reconstructions.

### 4. DISCUSSION

In this work, the quantitative capabilities of a clinical PET/CT device using \(^{44}\text{Sc}\) in a well-characterized phantom-based setup was investigated for the first time. Quantitative \(^{44}\text{Sc}\) PET/CT
Performances were compared with reference performance levels known for the well-established \(^{18}\text{F}\) PET imaging. We reported quantitative accuracy (BG calibration) and noise level (COV) assessed in a PVE-free background volume (the main volume of the NEMA/IEC NU2 phantom) and recovery coefficients obtained in spherical inserts of different size. In particular, we investigated the impact of different scatter models and PGC available in the vendor-based iterative reconstruction algorithm.

With regard to the noise, the results presented showed that, with clinically relevant activity concentrations, scan duration and reconstruction setups, noise levels below 15% can be achieved for a low number of iterations. These results, in terms of noise levels and, additionally, to the average residual lung error, are comparable to published results obtained with \(^{18}\text{F}\) FDG [31, 40] in similar clinical configurations for the same device.
Of note, it was found, according to the local dose calibrator cross-calibration, that the background quantitative signal was significantly below the acceptable performance levels considered for commonly-used PET radionuclides (for instance, BG$_{cal}$ out of the 1 ± 0.1 range defined as acceptable for $^{18}$F; Figure 1). In a recent publication [41], the authors reported the results from a survey across Dutch hospitals (8 centers, 13 different PET/CT), describing performance standard achievable in quantitative PET for both $^{68}$Ga and $^{18}$F. In particular, they found a system cross-calibration within 15% deviation in all tested devices and within 10% in 10 of 13 devices. We previously reported a background cross-calibration of <$5$% deviation using $^{18}$F for the mCT PET/CT system used in this work [31].

The low level of the measured background activity concentration could possibly be attributed to the specificity of the $^{44}$Sc decay (non-pure positron emission with associated high-energy prompt-gamma emission) which would significantly bias quantitative PET images in absence of optimized scatter and prompt-gamma corrections.

We initially considered the measured low level of BG calibration potentially ascribed to a non-appropriate scatter correction which, in the specific case, acted as an over-correction that resulted in an artificial depletion of the signal from the background volume. Further reconstructions were, therefore, performed using the scatter correction methods (relative and absolute) available from the vendor-based reconstruction algorithm implemented in the tested commercial PET/CT device.

Based on the results presented in this study, the available PGC improved the absolute quantitative accuracy of $^{44}$Sc PET, in agreement with results reported by Armstrong et al. [28] and Lubberink and Herzog [29] for other PET radionuclides. The impact of this correction is visible in Figure 2, where the overall scatter contribution was reduced, subsequently, improving the quantitative accuracy of the recovered activity concentration in the phantom background (Figure 1). The PGC fraction, although small, was not negligible: the difference between PGC reconstruction and non-PGC reconstructions was significant in terms of background calibration (p-values < 0.01 for non-PGC absolute and relative scatter corrections, respectively). Although the application of the PGC resulted in an improved quantitative accuracy, this was still well below accuracy levels obtained for $^{18}$F and $^{68}$Ga, which could indicate that further optimization could be performed. The average residual lung error was also found to be close to the reported $^{18}$F values for a similar device [40].

According to data presented in Figure 3, we observed that, even in the presence of a sub-optimal signal recovery, the uniform background, the signal recovery for spherical inserts with a diameter of <$20$ mm exceeded the reference range on values provided by the EANM/EARL recommendation [33]. It has been observed that RC values significantly higher than...
EANM/EARL reference values are commonly obtained when iterative reconstruction using both PSF and TOF information are used [34]. For this reason, the recommended range of RC values recommended by Kealep et al. was also considered (red shaded area in Figure 3). Recovery coefficients (RC_{A50}) were [31.5:66.8], [45.5:80.3], [66.4:75.6], [71.7:75.7], [75.1:78.6], and [76.7:80.9] for the 10, 13, 17, 22, 28, and 37 mm spherical inserts, respectively, on the reconstruction with absolute scatter and no PGC, confirming a quantitative signal recovery in the lower range of the values provided by Kealep et al. [34]. To compensate for the quantitative offset caused by the background calibration (BGcal < 1), we presented, for the optimal reconstruction setup (3 iterations), corrected RCs obtained by scaling up the measured RCs presented in Figure 3 by the respective BGcal offset compared to BGcal = 1 adopted in the NEMA approach. Corrected RCs are shown in Figure 4 and falls well within the range of values proposed by Kealep et al. [34], for PET reconstructions adopting TOF and PSF information. This observation further supports the hypothesis that achievable signal recovery in $^{44}$Sc PET is comparable to that achievable with $^{18}$F and $^{68}$Ga.

4.1. Limitations

This paper focused on the quantitative accuracy performances of a clinical PET scanner to quantify $^{44}$Sc. The present study does not investigate possible sources of quantitative bias, such as the effect of the positron range, in addition to the configured branching factor and its impact on quantification of $^{44}$Sc.

A possible explanation for the measured low level of background calibration could arise from the methodology adopted to determine the $^{44}$Sc activity. In this work the cross calibration relied on a reference activity value provided by PSI. The obtained factor was then used for all dispensing syringe activities. A different approach described in the literature [12, 42] defines the absolute $^{44}$Sc activity as the measured activity using the $^{18}$F dose calibrator settings multiplied by a factor (0.7). Although this approach has been tested, we found that this doesn’t correlate to our findings. By using this approach, we would have received 980.7 MBq at the reference time although the dispatched activity was 595.0 MBq. As a result, this approach was not further considered.

Phantom preparation and data analysis applied in our study differs in some points from the EARL (or NEMA) procedures (for instance, the lesion-to-background activity concentration ratio) and might impair a full comparability of results obtained with the cited procedures. Nevertheless, we consider that comparing our results with EARL proposed reference values (for the COV and RCs) would still be meaningful.

5. CONCLUSION

In this work, we evaluated the capability of a commercial PET/CT device to produce accurate quantitative imaging with $^{44}$Sc, for the first time. We performed dedicated $^{44}$Sc PET acquisitions using a controlled and clinical relevant phantom-based setup. Quantitative $^{44}$Sc PET performances, in terms of quantitative accuracy and noise levels, were shown to be comparable to that achieved in $^{18}$F PET. In particular, this study outlined that available scatter and prompt-gamma corrections plays an important role in $^{44}$Sc PET quantification. In view of a broader clinical application of $^{44}$Sc-based radiopharmaceuticals, it is essential to further improve and optimize these features.

DATA AVAILABILITY STATEMENT

The datasets generated for this study are available on request to the corresponding author.

AUTHOR CONTRIBUTIONS

All listed authors made substantial contributions in all four of the following categories.

- Substantial contributions to the conception or design of the work, or the acquisition, analysis, or interpretation of data for the work.
- Drafting the work or revising it critically for important intellectual content.
- Provide approval for publication of the content.
- Agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. More specifically, TL, SG, EN, CM, and NM coordinated the project and setup the framework. NM and CM coordinated the $^{44}$Sc production. TL and SG performed the phantom preparation, measurements, and analysis. PO contributed to the data analysis.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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