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PAPER

Positron range-free and multi-isotope tomography of positron emitters

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Abstract

Despite improvements in small animal PET instruments, many tracers cannot be imaged at sufficiently high resolutions due to positron range, while multi-tracer PET is hampered by the fact that all annihilation photons have equal energies. Here we realize multi-isotope and sub-mm resolution PET of isotopes with several mm positron range by utilizing prompt gamma photons that are commonly neglected. A PET-SPECT-CT scanner (VECTor/CT, MILabs, The Netherlands) equipped with a high-energy cluster-pinhole collimator was used to image ¹²⁴I and a mix of ¹²⁴I and ¹⁸F in phantoms and mice. In addition to positrons (mean range 3.4 mm) ¹²⁴I emits large amounts of 603 keV prompt gammas that—aided by excellent energy discrimination of NaI—were selected to reconstruct ¹²⁴I images that are unaffected by positron range. Photons detected in the 511 keV window were used to reconstruct ¹⁸F images. Images were reconstructed iteratively using an energy dependent matrix for each isotope. Correction of ¹⁸F images for contamination with ¹²⁴I annihilation photons was performed by Monte Carlo based range modelling and scaling of the ¹²⁴I prompt gamma image before subtracting it from the ¹⁸F image. Additionally, prompt gamma imaging was tested for ⁸⁹Zr that emits very high-energy prompts (909 keV). In Derenzo resolution phantoms 0.75 mm rods were clearly discernable for ¹²⁴I, ⁸⁹Zr and for simultaneously acquired ¹²⁴I and ¹⁸F imaging. Image quantification in phantoms with reservoirs filled with both 124I and 18F showed excellent separation of isotopes and high quantitative accuracy. Mouse imaging showed uptake of ¹²⁴I in tiny thyroid parts and simultaneously injected ¹⁸F-NaF in bone structures. The ability to obtain PET images at sub-mm resolution both for isotopes with several mm positron range and for multi-isotope PET adds to many other unique capabilities of VECTor's clustered pinhole imaging, including simultaneous sub-mm PET-SPECT and theranostic high energy SPECT.

1. Introduction

Preclinical PET and SPECT scanners are important devices for basic and translational research. The mouse is the most commonly used experimental animal due to its high similarity with the human homolog, today's existence of many mature genetic manipulation techniques, ease of fast breading and availability of economical housing. Typically, clinical scanners have resolutions ranging from 3 to 6 mm for PET and 8-10 mm for SPECT. As most mouse organs are roughly an order of magnitude smaller than their human counterparts, sub-mm resolution is required to measure molecule concentrations in similar sub-structures in organs and tumors. Preclinical SPECT most often relies on pinhole collimation and nowadays some systems reach resolutions down to a quarter mm in vivo (Ivashchenko et al 2015). Like clinical PET, preclinical PET is commonly based on coincidence detection of photons resulting from annihilation of an emitted positron with an electron in neighboring tissue. Resolutions down to 0.8 mm have been reported in high-end commercial coincidence PET systems (Yang et al 2004, Miyaoka et al 2004, Rouze et al 2004, Tai et al 2005) but only for isotopes with very small positron ranges

Figure 1. (A) Cluster of pinholes versus traditional pinhole. (B) clustered multi-pinhole collimator. Reproduced from Goorden *et al* 2020. © 2020 Institute of Physics and Engineering in Medicine. All rights reserved. (C) VECTor⁶CT system. Reproduced with permission from MIlabs.

Table 1. Positron ranges, half-lives, gamma energies and abundancies for several relevant PET isotopes (e.g. from (Le Loirec and Champion 2007a, 2007b, 2007c, Laforest and Liu 2009)). All isotopes in this table already have medical applications (Andreyev and Celler; Conti and Eriksson 2016).

Isotope	Positron range (Mean)	Positron range (Max)	Half-life	Probability positron emission	Energy and probability gamma emission
⁶⁸ Ga	3.56 mm	10.3 mm	67.8 min	88.9%	1077 keV (3.2%)
⁷⁶ Br	2.47 mm	20.2 mm	16.2 h	54.8%	559 keV (74%)
					657 keV (15.9%)
⁸² Rb	7.49 mm	18.6 mm	1.3 min	95.4%	777 keV (15.1%)
⁸⁶ Y	2.51 mm	11.1 mm	14.7 h	31.9%	1077 keV (82.5%)
					627 keV (32.6%)
^{124}I	3.37 mm	11.7 mm	100.2 h	22.7%	603 keV (62.9%)
					1691 keV (11.2%)
⁸⁹ Zr	1.27mm	4.21 mm	78.4 h	22.7%	909 keV (99%)
³⁸ K	5.67 mm	15.3 mm	7.61 m	99%	2170 keV (99%)
⁴⁴ Sc	2.46 mm	7.36 mm	3.97 h	94.3%	1157 keV (100%)
^{52m} Mn	5.29 mm	14.5 mm	21.1 m	98%	1434 keV (98%)
⁶⁰ Cu	4.13 mm	21.0 mm	23.4 m	92.5%	826 keV (21%)
					1332 keV (88%)
					1792 keV (46%)
⁷² As	5.19 mm	18.2 mm	26 h	88%	833 keV (815)

like 18 F. For a recent research prototype small field-of-view coincidence PET system dedicated to mouse brain imaging, a 0.6 mm resolution was reported (Yang *et al* 2016).

Many image degrading effects inherent to coincidence PET which play a relatively small role in human imaging become disturbing in small animals. Important factors in this regard are detector blurring, including depth-of-interaction (DOI) effects, and detection of random and scattered photons (Goorden and Beekman 2010, Goorden *et al* 2013). To reduce their impact, high-performance PET requires very expensive detector technology which can limit its practical application. For some isotopes an even more important image degrading effect is the positron range (e.g. 3.4 mm mean/ 11.7 mm max for ¹²⁴I, see table 1) resulting in significant losses in resolution and quantitative accuracy. Like other blurring effects these can be partly recovered using advanced iterative methods, but this requires accurate knowledge of tissue boundaries, e.g. from precisely aligned and preprocessed MRI or CT data (Cal-Gonzalez *et al* 2015). One should note that the shape of the 3D positron range kernel is highly peaked with long tails, which results in better small hot lesion visibility than would be the case for a Gaussian-like blurring kernel with the same mean range but that this on the other hand can cause more challenging quantification issues.

Earlier we have launched a method for simultaneous and sub-mm PET-SPECT imaging based on clustered pinhole collimation, named Versatile Emission Computed Tomography (VECTor (Goorden *et al* 2013, Walker *et al* 2014)). Due to the use of (i) clustered pinhole—rather than electronic-collimation (figure 1(A and B)) and (ii) gamma detectors with good energy resolution (8% at 511 keV), several image degrading effects inherent to electronic collimation are dramatically reduced or eliminated; for example scatter contamination in the photopeak is much lower, while detector blur and DOI have far less impact on resolution because of strong pinhole image magnification. At the same time, highly degrading noise equivalent count-rate effects caused by randoms and coincidence losses that severely affect coincidence PET are eliminated with clustered pinhole PET. Therefore, despite the lower number of photons detected with VECTor, it can for many cases outperform

coincidence PET in terms of image resolution and image contrast to noise ratio, particularly in organ and tumor imaging (Walker *et al* 2014). Unique capabilities of VECTor (figure 1(C)) include (i) performing sub-mm resolution PET and SPECT simultaneously, (ii) sub-mm resolution imaging of therapeutic isotopes that emit high energy gammas such as ²¹³Bi (440 keV) (de Swart *et al* 2016) and ¹³¹I (364 keV) (van der Have *et al* 2016), and (iii) providing an ultra-high resolution nuclear imaging platform with adaptable resolution-sensitivity trade-off e.g. through use of exchangeable collimators with application-specific pinhole diameters and geometries. As a consequence, this technology is already in use for a wide range of applications (e.g. (Walker *et al* 2014, Miwa *et al* 2015, de Swart *et al* 2016, van der Have *et al* 2016, Adachi *et al* 2017, Esquinas *et al* 2017, Robertson *et al* 2017, Chacko *et al* 2017, Verhoog *et al* 2018, Chekol *et al* 2018, Knight *et al* 2019a, Knight *et al* 2019b, Wilson *et al* 2019)).

Thus far, the above-mentioned positron range effect degrades VECTor's resolution as it does in coincidence PET. With VECTor often an equivalent SPECT isotope can be used as a work around (like ⁶⁷Ga instead of ⁶⁸Ga, ¹²³I, ¹²⁵I or ¹³¹I instead of ¹²⁴I, and ¹¹¹In instead of ⁸⁹Zr), isotopes which can all be imaged at sub-half-mm resolution. Here an additional option is presented in case one wants or needs to stick to PET isotopes, allowing for high-resolution imaging of long positron range isotopes. This option is enabled by VECTor's unique collimation technique and relies on the fact that several PET isotopes with long positron ranges also emit significant amounts of gammas straight from the atom (see table 1 showing that there are quite a number of such isotopes).

Another unmet need is to routinely create multi-isotope PET images. For coincidence PET systems thus far, two main methods have been proposed. A first method uses the difference in half-lives and kinetic behavior of different tracers, sometimes combined with staggered injection to separate their time-activity-curves (Rust et al 2006, Kadrmas et al 2013, Verhaeghe and Reader 2013). This has been applied in animal models (Black et al 2008, Figueiras et al 2011, Cheng et al 2015, Bell et al 2017) and patients (Joshi et al 2009, Zhang et al 2016). It relies on many assumptions about the pharmacokinetics of radiotracers or their spatial distribution and it has been pointed out that this is not actual simultaneous multi-tracer imaging (Fukuchi et al 2017) such as is for example done with SPECT systems that discriminate gammas emitted by different isotopes based on their energies. A second method is simultaneous imaging of a pure positron emitter and a positron emitter coemitting prompt gammas (Andreyev and Celler 2011, Gonzalez et al 2011, Andreyev et al 2014). The development of a small-animal multi-isotope PET based on this principle was reported recently (Fukuchi et al 2017). This method has the disadvantage that it requires modifications to the scanner's electronics to detect three gammas in coincidence and that a large number of quite bulky additional gamma detectors need to be added to reach a reasonable sensitivity for the scarce triple coincidences. In addition, the method works only for those PET isotopes that emit a high energy gamma *simultaneously* with a positron; it is therefore unsuited for delayed gammas or in cases when a large number of emitted gammas is not associated with positron decay but with electron capture.

The aim of the present paper is to (i) describe initial results of VECTor's capabilities of imaging isotopes with several mm positron range at sub-mm resolution and (ii) demonstrate simultaneous sub-mm imaging of different PET isotopes. Both new imaging capabilities are based on utilizing high-energy prompt or delayed gammas. Note that while VECTor uses gammas directly emitted by the isotope, it does not require triple coincidence and thus both delayed as well as prompt gammas not associated with positron emission can be utilized. In this paper we demonstrate these capabilities for the PET isotopes ⁸⁹Zr and ¹²⁴I in combination with ¹⁸F. Thus, positron range-free PET and dual-isotope PET are achieved by acquiring magnified multi-pinhole projections in the extremely high energy domain with a commercially available PET-SPECT-CT scanner.

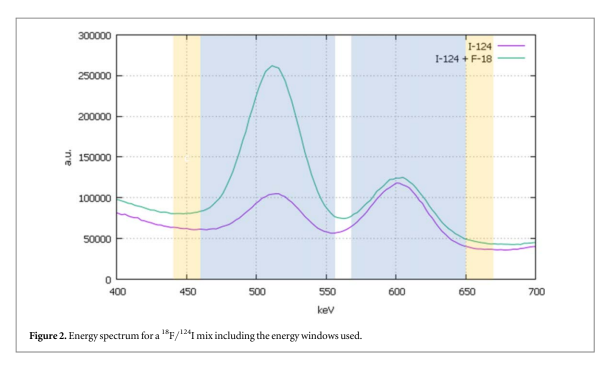
2. Materials and methods

2.1. Data acquisition

Animal and phantom scans were performed using a PET-SPECT-CT scanner (VECTor 6 CT, MILabs B.V.) equipped with three stationary large NaI(Tl) gamma detectors (595 \times 472, 9.5 mm crystal thickness) and a dedicated high energy mouse collimator (HE-UHR-M) with 144 pinholes (0.7 mm diameter each) organized in clusters of four (Goorden *et al* 2013). Images were acquired as list mode data meaning that estimated energy of each detected photon was stored. The advantage of this capability is that energy windows can be selected retrospectively.

2.2. In vivo animal imaging

All animal studies were performed in accordance with the Dutch Law on Animal Experimentation and all protocols were approved by the Animal Research Committee of the University Medical Center Utrecht. Healthy C57BL/6 mice (20–25 g body weight) were injected i.v. in the tail vein with 12 MBq of 124 I-NaI and 118 MBq of



 18 F-NaF, 24 h and 115 min before the scan started respectively. The animals were anesthetized using 2% of isoflurane in air and the tracer distributions were imaged simultaneously for 60 min.

2.3. Phantom imaging

A Derenzo phantom, containing 6 sectors of rods with varying diameter (0.45, 0.50, 0.55, 0.75, 0.80 and 0.85 mm) was used for resolution measurements. Per sector the rod diameters were equal to the distance between them. The phantom was filled with an initial activity of 29.9 MBq of ¹²⁴I and 29.9 MBq of ¹⁸F and scanned continuously for ten half-lives of ¹⁸F. For each half-life time scan, the first 30 min were reconstructed to obtain a series of images with different amounts of ¹⁸F. To additionally emulate the measurement of lower doses of ¹²⁴I, the list mode data of the last frame with completely decayed ¹⁸F was reduced by a factor that equals the fraction of the real dose and the emulated dose by randomly removing events from the list mode data. Additionally, single isotope ¹²⁴I Derenzo images were acquired with 26 MBq ¹²⁴I and a scan time of 30 min.

A cylindrical phantom (diameter 22 mm) containing three tubes (500 μ l per tube, inner diameter 6.5 mm) was imaged to assess quantitative accuracy. In this scan of 30 min, one tube contained 0.98 MBq of ¹²⁴I, a second tube was filled with 10.1 MBq of ¹⁸F, and the remaining tube contained a mix of 0.98 MBq of ¹²⁴I and 10.1 MBq of ¹⁸F.

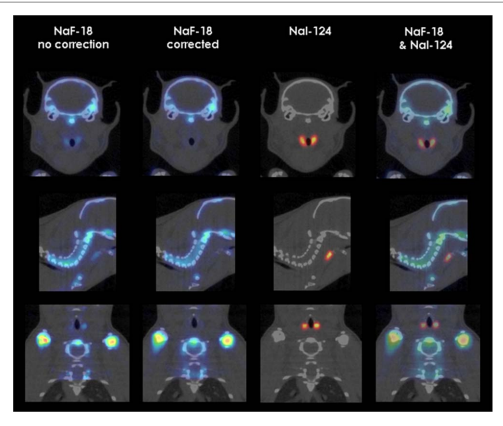
Images of a second Derenzo phantom (rod sizes of 0.70, 0.75, 0.80, 0.90, 1.00 and 1.20 mm) were acquired with an initial dose of 40 MBq 89 Zr and scan time of 30 min.

2.4. Image reconstruction

All images were reconstructed by a combined dual-matrix dual-voxel pixel-based (Branderhorst *et al* 2010) similarity regulated (Vaissier *et al* 2016) OSEM (DM-SR-OSEM) algorithm. Dual-matrix image reconstruction (Zeng and Gullberg 1997) uses different matrices for forward projection and back projection to accelerate reconstruction. In our case, the back projection step did not contain positron range blurring and included only part of the photons penetrating the collimator (for details see (Goorden *et al* 2020)). Dual-voxel reconstruction is an acceleration technique (Goorden *et al* 2020) that uses larger voxels for the low frequency tail part of the point spread functions in the forward projection. MC-generated detector PSFs including Compton scatter were used. A triple energy window scatter and cross talk correction (Ogawa *et al* 1991) was used and was modified for this specific case of dual-isotope imaging (see figure 2). For ¹⁸F imaging, photons in the 511 keV photopeak (461–561 keV) were used as well as 2 background subtraction windows (441–461 keV / 650–670 keV). For ¹²⁴I imaging, the 609 keV photopeak (570–650 keV) was used with one background window (650–670 keV). For ⁸⁹Zr imaging, reconstructions from (i) the 511 keV photopeak (461–561 keV) and (ii) the 909 keV photopeak (841–977 keV) were compared. Scatter was corrected by using a triple energy window with a width of 2.5% each on each side of the 15% wide peak window.

Separate matrices were used to model the energy specific photon transport through collimator and detector material for 511, 603 or 909 keV photons. Matrices were obtained by raytracing to model the pinhole penetration and detector interaction for the specified energy, while calibration was based on ^{99m}Tc point source

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 $\textbf{Figure 3.} \ Simultaneous \ dual \ isotope \ mouse \ images \ with \ ^{18}F-NaF \ and \ ^{124}I-NaI \ (12\ MBq \ and 55\ MBq \ resp. \ at \ start) \ overlaid \ with \ x-ray \ CT \ images. Top: Transaxial slices. Center: Sagittal slices. Bottom: coronal slices. Left and 2nd column: \ ^{18}F-NaF \ image \ uncorrected \ and corrected for down scatter and annihilation photons originating from \ ^{124}I-NaI. Third Column: \ ^{124}I-NaI \ image \ from 603 \ keV \ prompt \ photons \ showing \ small \ details from the thyroid. Right Column: Corrected \ ^{18}F-NaF \ and \ ^{124}I-NaI \ image \ merged.$

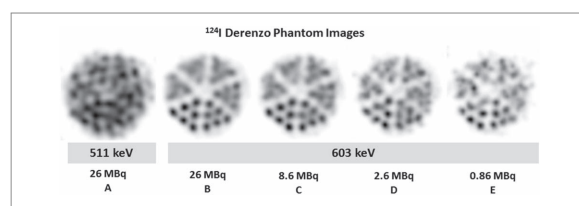


Figure 4. Images of Derenzo phantom (rod sizes 0.45, 0.50, 0.55, 0.75, 0.80 and 0.85 mm) filled with different amounts of ¹²⁴I and scanned for 30 min. (A) Reconstruction from 511 keV annihilation photons. (B)–(E) Prompt images at different activity levels (up to a factor of 30 difference with frames A and B).

measurements (Goorden *et al* 2016). For absolute quantitative imaging a calibration method based on a single cup of activity was used. This was applied to each isotope with single isotope imaging (Wu *et al* 2011).

2.5. Calibration for cross talk correction

For 124 I imaging only the 603 keV prompt gammas were selected, while the 511 keV photons were not used because of the enormous positron range. When 18 F is imaged simultaneously with 124 I using the 511 keV channel, one in fact creates images that represent positron annihilations of both isotopes where the 124 I image is heavily blurred by positron range effects. To correct the 18 F image for this effect we calculated an estimate of the amount of contamination of 124 I present in the 18 F image with the following method. First positron range blurring was applied to the positron range-free 603 keV 124 I image using pre-calculated and normalized Gate Monte Carlo simulation (Jan *et al* 2004) generated kernels. This blurred image was subsequently scaled and subtracted from a 511 keV 124 I only image determined experimentally from phantom experiments. The scaling

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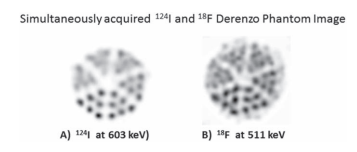


Figure 5. Derenzo phantom (rod sizes 0.45, 0.50, 0.55, 0.75, 0.80 and 0.85 mm), activity ratio ^{124}I : ^{18}F equals 1:1. (A) ^{124}I image. (B) 511 keV ^{18}F image corrected for ^{124}I annihilation photons.

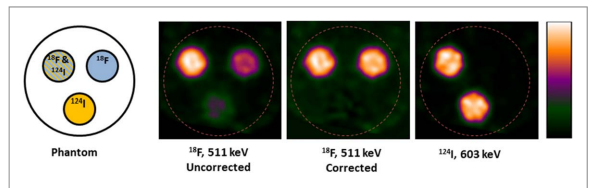


Figure 6. Quantification phantom with 3 compartments filled according to the concentrations provided in table 2. This table also shows that the quantitative accuracy for 124 I is excellent and that after correction for 124 I cross talk the same holds true for 18 F.

factor was considered to be optimal when after subtraction there are zero counts left. The scaling factor found this way was 0.87. Using this scaling factor 18 F images obtained from dual-isotope 18 F/ 124 I imaging were corrected by subtracting the estimated contamination of 124 I.

3. Results

Figure 3 shows images of the mouse co-injected with 124 I-NaI and 18 F-NaF. The activities at the time of imaging were 12 MBq and 55 MBq respectively. By only using the prompt gammas for 124 I-NaI reconstruction, the 3.4 mm average positron range of 124 I can be avoided and structures smaller than a mm in the mouse thyroid can be easily resolved. Correction of 18 F-NaF images is necessary to remove contamination from the 124 I-NaI distribution from the images. Images were reconstructed using 50 iterations DM-SR-OSEM and 3D Gaussian post-filtered with FWHM = 0.6 mm.

Figure 4 shows the difference between 124 I reconstructions from 511 keV annihilation photon imaging (Frame A) and 603 keV photons (Frame B) at equal dose (images were reconstructed from the same scan). Frames B–E show the effect of count reduction corresponding to a range of activities between 26 MBq and 0.86 MBq (at the start of the scan). All images were reconstructed using 50 iterations DM-SR-OSEM and Gaussian post-filtered with a FWHM of 0.5 mm, 0.5 mm, 0.55 mm, 0.60 mm and 0.65 mm respectively to get proper visualization at the increasing noise level with lower activity.

Figure 5 shows simultaneous Dual-Isotope PET images of a Derenzo phantom filled with a mix of 124 I and 18 F discerning the 0.75 mm rods. Images were reconstructed using 50 iterations DM-SR-OSEM and Gaussian post-filtered (FWHM = 0.5 mm). In the 511 keV photopeak the number of counts after background correction amounted to 72.0 M, which is the estimated number of primary counts from 18 F, and in the 603 keV window the number of counts after background correction amounted to 30.5 M.

Figure 6 shows the phantom with 3 compartments with (1) a 18 F/ 124 I mix of activity concentration ratio of 1:3.86, (2) 18 F only and (3) 124 I only. The amounts of activity were calculated from the images using cylindrical VOIs of 8 mm diameter and 30 mm length around the cups. Images were reconstructed using 50 iterations DM-SR-OSEM and 3D Gaussian post-filtered with an FWHM of 1 mm. Reconstructed amounts of 18 F in compartments (1) and (2) were found to be equal as was the amount of 124 I in compartments (1) and (3). The intensity images together with the ROI values in table 2 show that after correction, the quantitative accuracy is excellent. The activity in the filled compartments barely deviates from the true concentrations but for 18 F, application of cross talk correction is very important.

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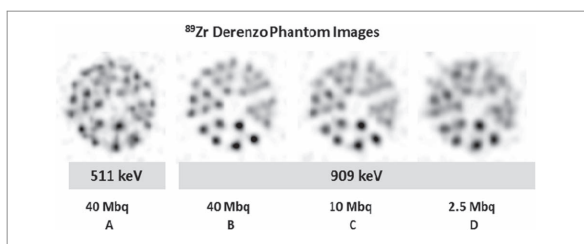


Figure 7. Derenzo phantom (rod diameters $1.2\,1.0\,0.9\,0.8\,0.75\,0.7\,\text{mm}$) filled with $^{89}\text{Zr.}$ (A) image based on the use of 511 keV window, compared to (B) image from the 909 keV photopeak window. (C)–(D) Prompt images at lower activity levels (up till $16\times$ lower than in frame A).

Table 2. Quantitative accuracy in cup phantom shown in figure 6 with ¹²⁴I only, ¹⁸F only or a mix of both.

	$\begin{array}{c} \text{Measured Concentration (MBq ml}^{-1}) \\ \text{Uncorrected} \\ \text{\% Error} \end{array}$	$\begin{tabular}{ll} Measured Concentration (MBq ml$^{-1}$) \\ Corrected \\ \% Error \end{tabular}$	True Concentration (MBq ml^{-1})
Compartment I			
¹⁸ F	0.54 (54%)	0.34(3%)	0.35
^{124}I	1.42 (5%)	n.a.	1.35
Compartment II			
¹⁸ F	0.36 (3%)	0.34(3%)	0.35
^{124}I	0.05 (4%)	n.a.	0.0
Compartment III			
¹⁸ F	0.21 (60%)	0.01 (3%)	0.0
^{124}I	1.42(5%)	n.a.	1.35

Figure 7 shows that also ⁸⁹Zr images can strongly improve in clarity when prompts are imaged instead of annihilation photons, despite the fact that the photon energies are almost 1 MeV. All images were reconstructed using 100 iterations DM-SR-OSEM and Gaussian post-filtered with a FWHM of 0.5 mm, 0.5 mm, 0.6 mm and 0.8 mm respectively to get proper visualization at the increasing noise level with lower activity. At 511 keV many rods are affected in shape by scatter and positron range although even 0.7 mm rods are visible, but 0.75 mm rods are not well discernable from each other. The rods at 909 keV are much rounder and therefore more realistic, but the 0.7 mm rods are hard to see at 909 keV which may be explained by a wider PSF at 909 keV due to e.g. pinhole edge penetration. It should be noted that scatter and the amount of counts play a big role in the quality of the 511 keV based images: for the 40 MBq scan in the 511 keV window the photopeak counts amounted 173 M but the estimated number of primary counts was 34.1 M after TEW correction. This is 3.35 times lower than the amount of estimated primary photons (again by TEW) of 114.5 M counts in the 909 keV window.

4. Discussion

We presented first results of positron range-free PET imaging based on imaging prompt gammas that are coemitted with positrons by many PET isotopes and often have a considerable abundance (table 1). We tested this method for ¹²⁴I and ⁸⁹Zr and showed that for both isotopes 0.75 mm rods in a Derenzo resolution phantom could be clearly discerned (figures 4 and 7), which used to be impossible before. Since partial volume effects are a main hurdle for quantitative imaging, the presented method can be a great step forward to precise imaging of these isotopes. Since the presented work was carried out with an isotope with a large positron range (3.4 mm in average) for coincidence imaging, and one with an extremely high prompt energy (909 keV), we expect that many other applications with isotopes listed in table 1 can benefit from imaging the prompt gammas as well.

Also combining the information from all photopeaks (511 keV plus prompt gammas) may be interesting to further improve the results in some studies.

The current work was carried out with a VECTor system with rather thin crystals. Today these systems are also delivered with crystals that have more than $2.3 \times \text{higher}$ capturing efficiency and are equipped with dedicated software that can model the increased DOI effects including Compton effects in the crystal. Such a scanner was not available to us in an animal lab at this stage of the research project. In addition, higher sensitivity collimators can be used which together would support much shorter scan times or lower doses than used in the present paper, albeit at the cost of some spatial resolution The loss of resolution can be partly compensated for by accurate modeling of the collimator in the system matrix that is used for reconstruction.

So far energy windows settings and image reconstruction have not yet been optimized. The optimal width of the windows may be dose dependent and using both the 511 keV window and window for prompt photons could be interesting when for each window the reconstruction matrix is optimized.

Simultaneous dual-isotope PET imaging is important since it can reduce scan time compared to two separate scans which can limit the time needed to keep the animal under anesthesia and it inherently results in perfectly registered images of different tracer molecules. The present work shows that such studies are feasible even based on simple cross talk correction.

5. Conclusion

This study showed that (i) sub-mm resolution imaging of a PET isotope with several mm mean positron range and (ii) simultaneous sub-mm resolution imaging of different PET isotopes is enabled by clustered pinhole collimation and magnification, using a system with stationary large field-of-view NaI detectors (VECTor) and dedicated image reconstruction methods. Many other PET isotopes with a large positron range also have also additional prompt gammas that can be imaged in this way. The use of a wide variety of PET tracers, including large positron-range PET isotopes, and mixes of multiple PET and SPECT tracers paves the way for many new imaging protocols in biomedical research.

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