

Document Version

Final published version

Licence

CC BY

Citation (APA)

Goorden, M. C., Kamphuis, C., Ramakers, R. M., & Beekman, F. J. (2020). Accelerated image reconstruction by a combined dual-matrix dual-voxel approach. *Physics in Medicine and Biology*, 65(10), Article 105014. <https://doi.org/10.1088/1361-6560/ab82e9>

Important note

To cite this publication, please use the final published version (if applicable). Please check the document version above.

Copyright

In case the licence states "Dutch Copyright Act (Article 25fa)", this publication was made available Green Open Access via the TU Delft Institutional Repository pursuant to Dutch Copyright Act (Article 25fa, the Taverne amendment). This provision does not affect copyright ownership. Unless copyright is transferred by contract or statute, it remains with the copyright holder.

Sharing and reuse

Other than for strictly personal use, it is not permitted to download, forward or distribute the text or part of it, without the consent of the author(s) and/or copyright holder(s), unless the work is under an open content license such as Creative Commons.

Takedown policy

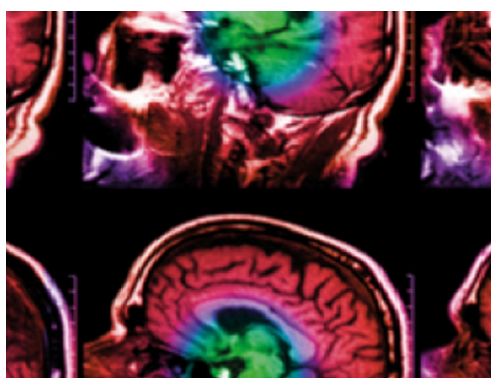
Please contact us and provide details if you believe this document breaches copyrights. We will remove access to the work immediately and investigate your claim.

PAPER • OPEN ACCESS

Accelerated image reconstruction by a combined dual-matrix dual-voxel approach

To cite this article: Marlies C Goorden *et al* 2020 *Phys. Med. Biol.* **65** 105014

View the [article online](#) for updates and enhancements.



IPEM | IOP

Series in Physics and Engineering in Medicine and Biology

Your publishing choice in medical physics,
biomedical engineering and related subjects.

Start exploring the collection—download the
first chapter of every title for free.



PAPER

OPEN ACCESS

RECEIVED
8 November 2019REVISED
6 March 2020ACCEPTED FOR PUBLICATION
24 March 2020PUBLISHED
29 May 2020

Original content from this work may be used under the terms of the [Creative Commons Attribution 3.0 licence](#).

Any further distribution of this work must maintain attribution to the author(s) and the title of the work, journal citation and DOI.



Accelerated image reconstruction by a combined dual-matrix dual-voxel approach

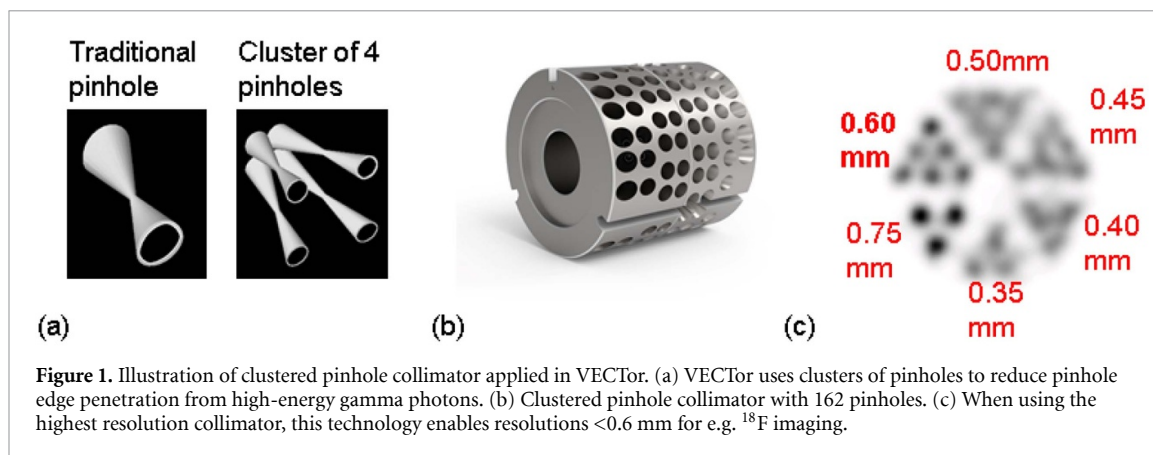
Marlies C Goorden^{1,4}, Chris Kamphuis^{2,3}, Ruud M Ramakers^{1,2,3} and Freek J Beekman^{1,2,3}¹ Section of Biomedical Imaging, Faculty of Applied Sciences, Delft University of Technology, Mekelweg 15, 2629 JB Delft, The Netherlands² MILabs B.V, Heidelberglaan 100, 3584 CX Utrecht, The Netherlands³ Department of Translational Neuroscience, Brain Center Rudolf Magnus, University Medical Center Utrecht, Utrecht, The Netherlands⁴ Author to whom any correspondence should be addressed.E-mail: m.c.goorden@tudelft.nl**Keywords:** image reconstruction, positron emission tomography (PET), single photon emission computed tomography (SPECT)

Abstract

Today, versatile emission computed tomography (VECTor) technology using dedicated high-energy collimation enables simultaneous positron emission tomography (PET) and single photon emission computed tomography (SPECT) down to 0.6 mm and 0.4 mm resolution in mice, respectively. We recently showed that for optimal resolution and quantitative accuracy of PET images the long tails of the 511 keV point spread functions (PSFs) need to be fully modelled during image reconstruction. This, however, leads to very time consuming reconstructions and thus significant acceleration in reconstruction speed is highly desirable. To this end we propose and validate a combined dual-matrix dual-voxel (DM-DV) approach: for the forward projection the slowly varying PSF tails are modelled on a three times rougher voxel grid than the central parts of the PSFs, while in the backprojection only parts of the PSF tails are included. DM-DV reconstruction is implemented in pixel-based ordered subsets expectation maximization (POSEM) and in a recently proposed accelerated pixel-based similarity-regulated ordered subsets expectation maximization (SROSEM). Both a visual assessment and a quantitative contrast-noise analysis confirm that images of a hot-rod phantom are practically identical when reconstructed with standard POSEM, DM-DV-POSEM or DM-DV-SROSEM. However, compared to POSEM, DM-DV-POSEM can reach the same contrast 5.0 times faster, while with DM-DV-SROSEM this acceleration factor increases to 11.5. Furthermore, mouse cardiac and bone images reconstructed with DM-DV-SROSEM are visually almost indistinguishable from POSEM reconstructed images but typically need an order of magnitude less reconstruction time.

1. Introduction

Today, small animal positron emission tomography (PET) and single photon emission computed tomography (SPECT) are key for discovering disease mechanisms, developing pharmaceuticals, radionuclide therapies and novel diagnostic radiotracers (Weissleder *et al* 2010). Preclinical SPECT commonly uses multi-pinhole collimation to image the distribution of single gamma emitting tracers and reaches uniform resolutions of 0.25 mm *in vivo* in the entire mouse (Ivashchenko *et al* 2015) and 0.12 mm *ex vivo* using a tissue sample collimator (Nguyen *et al* 2019). In contrast, the majority of PET scanners are based on electronically collimated coincidence imaging of 511 keV annihilation photon pairs. At the moment the best commercial coincidence PET scanners have a resolution down to 0.9 mm (Lecomte *et al* 1994, Cherry *et al* 1997, Ziemons *et al* 2005, Surti *et al* 2005, Visser *et al* 2009, Zhang *et al* 2011, Szanda *et al* 2011, Sanchez *et al* 2012, Herrmann *et al* 2013, Nagy *et al* 2013, Sato *et al* 2016, Krishnamoorthy *et al* 2018), while a prototype coincidence PET scanner with a central resolution of 0.6 mm has been reported (Yang *et al* 2016). We have taken a different approach by developing versatile emission computed tomography (VECTor) (MILabs BV,



(Beekman 2011, Goorden *et al* 2013)), a preclinical scanner based on clustered pinhole technology (Goorden and Beekman 2010) that is suitable for imaging gammas in the 27 keV up to 1 MeV energy range (including 511 keV annihilation photons) at high resolutions (<0.6 mm for ^{18}F -PET, see figure 1, and <0.75 mm for ^{124}I -PET). Thus PET tracer imaging with VECTor is not based on coincidence detection of pairs of photons but instead annihilation photons are imaged as singles. VECTor can image PET tracers simultaneously with one or multiple SPECT tracers based on energy discrimination of detected photons. In some cases it is even possible to perform sub-mm resolution multi-isotope PET, using the fact that many PET tracers also emit high-energy prompt gammas with different energy than the annihilation photons (Beekman *et al* 2018). Simultaneous acquisition of multiple isotope tracer distributions under identical conditions can be crucial e.g. when the physiological functions studied depend on each other or when different tracers have to be compared 1:1. The latter could have applications in tracer research if one would like to compare uptake of different tracers targeting the same biological function in the same animal at the same time. In addition, VECTor has applications in the emerging domain of theranostic tracers which often emit high-energy gammas, e.g. for ^{213}Bi (440 keV, (de Swart *et al* 2016) or ^{131}I (364 keV, (van der Have *et al* 2016)). For these reasons VECTor is in use and has been evaluated in many biomedical labs worldwide (Walker *et al* 2014, Ter Heine *et al* 2014, Lange *et al* 2015, Miwa *et al* 2015, Vangestel *et al* 2016, Dalm *et al* 2017, Adachi *et al* 2017; Esquinas *et al* 2017, Dogan *et al* 2017, Robertson *et al* 2017, Chacko *et al* 2017, Chekol *et al* 2018, Verhoog *et al* 2018, Kim *et al* 2019).

Key to accurate image reconstruction is the use of statistical iterative image reconstruction algorithms (Qi and Leahy 2006). In order to maximize resolution and quantitative accuracy, these algorithms ideally use accurate models for transport of gamma photons through tissue, collimator material and in the detector which can be represented by energy-dependent point spread functions (PSFs) that together form the system matrix. Each element in the system matrix represents the probability that a gamma photon emitted from a certain voxel ends up being detected in a certain detector pixel. In (Goorden *et al* 2016) we presented ray tracing software to efficiently calculate photon transport of 511 keV annihilation photons through collimator structures and in the gamma detector. We found that this software enabled to obtain accurately reconstructed images of ^{18}F provided that also large parts of the PSF tails are modelled. These tails contain gamma photons that pass through large amounts of collimator material and have therefore a low probability of not being attenuated. We showed that including photons with a probability as low as even 1% of passing through the collimator contributes to significantly improved contrast-noise characteristics. For imaging of conventional SPECT tracers (140 keV energy) including such low probability gamma photon paths does not improve images significantly (van der Have *et al* 2008). In (Goorden *et al* 2016) we presented a probable explanation for this discrepancy; we calculated that at photon energies of 140 keV, 83% of the photon flux was contained in photon paths with a probability $>20\%$ to pass through the collimator (in case of a large uniform activity distribution). However, at photon energies of 511 keV, only 35% of the photon flux was modelled by these high-probability photon paths. Thus 511 keV photon transport is not modelled accurately in reconstruction if low-probability photon paths are not included as they contribute significantly to the amount of detected photons.

Unfortunately, including a relatively high fraction of low-probability gamma photon paths has its disadvantage in terms of reconstruction speed. In this paper we propose a new dual-voxel (DV) approach to tackle this issue. It is based on the general observation that the PSF tails are much more gradually decreasing in intensity than the central PSF part. As a consequence, neighboring voxels in image space that project into the PSF's tails will have matrix elements that are very close in value. Thus, using a single mean matrix

element for such neighboring voxels may still enable to calculate an accurate projection. This boils down to using larger voxels for that part of the system matrix that contains slowly varying components while for quickly varying components a smaller voxel size is used. In this paper we will combine this DV approach with a dual-matrix (DM) approach (Zeng and Gullberg 1992, 2000, Welch and Gullberg 1997, Kamphuis *et al* 1998, Kadrmas *et al* 1998, Beekman *et al* 2002) in which a much smaller system matrix is used for the backprojection step of image reconstruction than for forward projection. We integrated this DM-DV approach in a pixel-based ordered subsets expectation maximization (POSEM) (Branderhorst *et al* 2010) algorithm as well as in a pixel-based similarity-regulated OSEM (SROSEM) algorithm. The latter was recently developed in our group to accelerate image reconstruction by safely allowing the use of more subsets than with standard POSEM (Vaissier *et al* 2016). We apply DM-DV reconstruction to phantom images and animal scans to investigate how much image reconstruction can be accelerated without loss of image quality.

2. Materials and methods

2.1. VECTor PET-SPECT system

The mouse VECTor collimator (figure 1) used here contains 162 clustered pinholes, each 0.75 mm in diameter. The system used was a VECTor/CT system (MILabs B.V. Utrecht, The Netherlands) that applies a triangular detector set-up with three PMT-based large-area NaI(Tl) gamma detectors. The maximum count yield of a clustered pinhole collimator is in the central field of view (FOV) and therefore a special graphical user interface is used that allows to focus on parts of the body (Branderhorst *et al* 2011). Larger regions up to whole body scans are done by stepping the animal through the central focus volume (Vastenhouw and Beekman 2007). In this paper a spiral bed sequence was used for data acquisition (Vaissier *et al* 2012). More details on collimator and scanner design can be found in (Goorden *et al* 2013).

2.2. System matrix calculation

The system matrix for 511 keV gamma photons is calculated as described in (Goorden *et al* 2016). In this procedure a scanning ^{99m}Tc point source measurement is used for calibration. From the point source measurement a number of geometrical parameters for VECTor's geometry are estimated; these parameters describe a translation and rotation of collimator tube and of the three gamma detectors with respect to the designed positions and orientations. After the geometry has thus been determined, a voxelized raytracing simulation is employed to calculate system matrix elements. The voxelized raytracer was developed in-house and is described in detail in (Wang *et al* 2017). It uses a voxelized model of collimator and detector (0.1 mm cubic voxels in this case). To each of the voxels an attenuation coefficient is assigned and photon transport through the voxelized volume is calculated. In this simulation, gamma photon attenuation in the collimator is included, but scatter is ignored. Variable depth-of-interaction in the gamma detector is also included by modelling the varying interaction position in the scintillation crystal of the gamma detector using Beer's law. Additional detector non-idealities are incorporated by applying Gaussian detector blurring with 3.5 mm full-width-at-half-maximum (FWHM). Note that this system matrix calculation is different from the method described in (van der Have *et al* 2008) to obtain the SPECT system matrix, although the same point source measurement is used.

Storing system matrix elements for all possible voxel-detector pixel combinations is not feasible as this would result in about 10^{12} system matrix elements for VECTor for a voxel size of 0.4 mm (the typical choice for high-resolution PET images). Therefore, we implemented a cut-off C in the raytracing code with a value between 0 and 1. Only gamma photon paths with a probability $< C \cdot 100\%$ to pass through the collimator are incorporated into the system matrix. In (Goorden *et al* 2016) it was shown that the above-described method for calculating the system matrix yields accurate high-resolution images if a cut off as low as 0.01 is used. Therefore, throughout the main part of the paper a value of C of 0.01 was applied (while in the appendix some results for C of 0.05 are provided as well).

2.3. Image reconstruction

In this work different image reconstruction algorithms are tested and compared in terms of speed and quality of reconstructed images. In all cases, the positron range was only included in the forward projection (which effectively results in a DM approach (Zeng and Gullberg 1992, Kamphuis *et al* 1998)). Photo peak windows were set to a width of 20% of the photo peak value. Scatter correction was performed using a triple energy window correction (King *et al* 1997) with side windows adjacent to the photo peak having a width of 80 keV. Reconstructions were performed both at 0.4 mm and 0.8 mm voxel size, as both sizes are used in practice. While for both voxel sizes reconstruction times are compared, all displayed images in this paper have 0.4 mm voxel size.

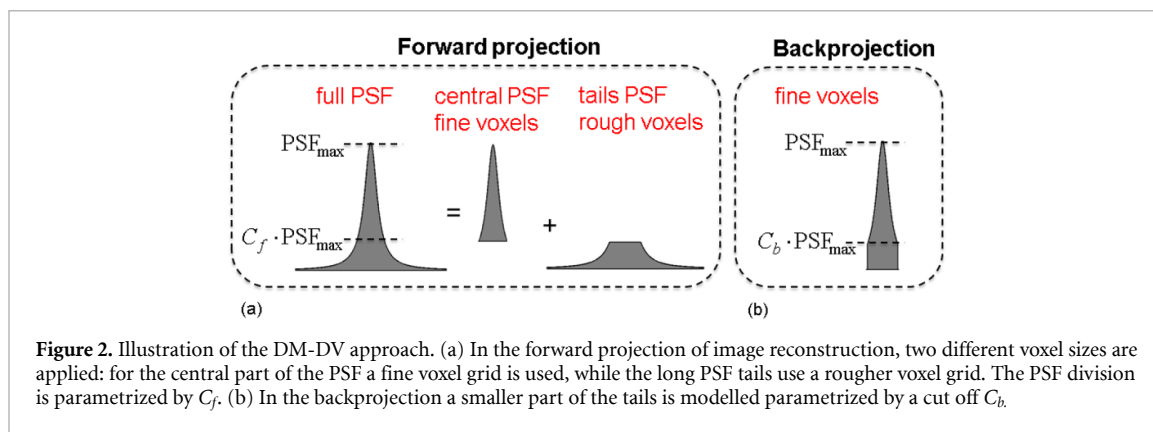


Figure 2. Illustration of the DM-DV approach. (a) In the forward projection of image reconstruction, two different voxel sizes are applied: for the central part of the PSF a fine voxel grid is used, while the long PSF tails use a rougher voxel grid. The PSF division is parametrized by C_f . (b) In the backprojection a smaller part of the tails is modelled parametrized by a cut off C_b .

As the gold standard reconstruction algorithm, we use POSEM (Hudson and Larkin 1994, Branderhorst *et al* 2010) with 32 subsets as this is the reconstruction method that was tested and widely applied in our previous works (Goorden *et al* 2013, 2016). POSEM is an ordered subset algorithm with pixel-based subset choice which leads to a better subset balance than the projection-view-based choice in standard OSEM and therefore allows for much larger acceleration factors over the maximum likelihood estimation method (MLEM) (Branderhorst *et al* 2010). Secondly, pixel-based SROSEM (Vaissier *et al* 2016) was tested. SROSEM is an accelerated image reconstruction algorithm that we recently proposed and that automatically and locally adapts the number of subsets used based on the similarity of update terms for different subsets. This way pixel-based SROSEM can safely use a much larger number of subsets than POSEM for high-count regions and therefore further accelerate image reconstruction. In (Vaissier *et al* 2016) SROSEM was validated for multi-pinhole SPECT, here we apply the algorithm to PET tracer imaging with VECTOR. We set a maximum number of subsets of 128 and a similarity threshold value of 40%.

To further accelerate image reconstruction, we implemented a DV approach to accelerate the forward projection step while a DM approach was used to accelerate the backprojection. The DV approach is illustrated in figure 2(a). In this figure a PSF is shown with its maximum value denoted by PSF_{max} . It can be seen in this image that the central part of the PSF is changing more rapidly than the tails of the PSF which are more slowly varying. The idea behind the DV approach we use is that in order to calculate an accurate forward projection, it may not be necessary to represent the tails of the PSF on the same fine voxel grid as the central part. To this end, a tail cut-off parameter C_f ($0 < C_f < 1$) is defined to divide the PSF into two parts; a ‘central PSF’ and a ‘tails PSF’. Based on the fact that the central part contains the highest spatial frequencies while the tails are more slowly varying, we set a three times coarser voxel size for the tails PSF. We thus assume that the ‘tails PSF’ corresponding to neighboring voxels is quite similar and that the simulated forward projection is not very different if one uses the ‘tails PSF’ of a nearby voxel in the forward projection. During image reconstruction the forward projection is then assumed to be the sum of (i) the forward projection obtained by letting the central PSF act on the fine voxel activity distribution and, (ii) the forward projection of the tails PSF which acts on a activity distribution binned on the larger voxel size.

Secondly, for the backprojection step we use a matrix with a higher tail cut off C_b than in the forward projection. Such a DM approach in which full photon transport is not taken into account in the backprojection step has been used previously and it has been shown to often lead to the same quality images as a standard reconstruction (Zeng and Gullberg 1992, Kamphuis *et al* 1998). Thus in the backprojection step of our algorithm, tails were not included.

We integrated the DM-DV approach both in the POSEM and the SROSEM algorithm with settings as described above. Furthermore, in this work we fixed parameters C_f and C_b to 0.05. This value of 0.05 is based on preliminary try-outs in our lab but not on a rigorous optimization study. Note that these cut off parameters are significantly larger than the value of the cutoff parameter $C = 0.01$ of the whole PSF (section 2.2).

Matrices with higher cut offs were obtained from the matrix with lowest cut off in the following way. First each matrix element in the lower cut off matrix ($C = 0.01$) was divided by the detector efficiency and by a geometrical factor that quantifies the probability of the gamma photon reaching that particular detector pixel if no collimator would be present. This division is done as both these factors were incorporated in the system matrix elements (see (Goorden *et al* 2016) for more details). The remaining value after division is a measure of the probability of transmission through the collimator and if it exceeds the higher cutoff value (i.e. 0.05) the matrix element is saved to the higher cut off matrix.

To obtain the tails and central PSF matrices we follow the following procedure. Let the original matrix element (including detector efficiency and geometrical factor) have a value of M while the remaining matrix element after division is m . If $m > C_f$, then a matrix element of value $\left(1 - \frac{C_f}{m}\right) \cdot M$ is saved to the central PSF matrix, while the tails PSF matrix is considered to have a value of $\frac{C_f}{m} \cdot M$ (as also illustrated in figure 2(a)). On the other hand, if $m < C_f$, the matrix element M is solely saved to the tails PSF matrix. However, an element is only saved to the tails PSF matrix if the fine voxel corresponding to this matrix element is the central voxel on the rougher voxel grid of the tails PSF matrix (so only for 1 out of 27 fine voxels do we save a matrix element to the tails matrix).

Note that the procedure is the same for every matrix element and the exact shape of the PSF is not taken into account, i.e. there is no special procedure for PSFs near the edge of the FOV which may have a very asymmetrical appearance. In the [appendix](#) of this paper (figure A1) we provide a few sample PSFs for pairs of neighboring voxels, one pair near the centre of the pinhole's FOV and another pair near its edge to get an idea of their appearance.

With this procedure we found that while matrices with a cut off of 1% contained on average $20 \cdot 10^9$ matrix elements, the 5% cut off matrix used in backprojection contained on average $4.2 \cdot 10^9$ elements and for the tail matrix this number decreased to $0.75 \cdot 10^9$ elements. The average is over different VECTor systems which may have slightly different system matrices.

2.4. Phantom studies

2.4.1. Hot-rod Derenzo phantom.

A hot-rod Derenzo resolution phantom was filled with a 54 MBq ^{18}F solution and scanned for 4 h. The phantom had six sectors each containing a set of equally sized capillaries with diameters of 0.7, 0.8, 0.9, 1.0, 1.2 and 1.5 mm. The distance between capillary centres was equal to twice the capillary diameter. For visualization a 0.5 mm FWHM 3D Gaussian filter was applied to reconstructed images.

2.4.2. Uniformly filled syringe.

A 12 ml syringe (15.9 mm inner diameter, 6 cm length) was filled with 65 MBq ^{18}F solution and scanned for 3.5 h. For visualization, images were filtered with a 0.8 mm FWHM 3D Gaussian.

2.5. Animal studies

Animal studies were carried out in accordance with the Dutch Law on Animal Experimentation and conducted according to protocols approved by the Animal Research Committee of the University Medical Center Utrecht. In order to judge better if even small deviations between applied algorithms appear, high count scans with relatively long acquisition times were selected for the present paper. These studies were previously used in our paper describing the POSEM reconstruction software (Goorden *et al* 2016).

2.5.1. Mouse cardiac scan.

A 29 g C57Bl6 mouse was anesthetized with isoflurane and subsequently injected intravenously with 24 MBq ^{18}F -deoxyglucose (^{18}F -FDG). A 30 min acquisition began 10 min post injection. For visualization, images were post-filtered with a 0.8 mm FWHM 3D Gaussian.

2.5.2. Mouse bone scan.

A 33 g C57Bl6 mouse was anesthetized with isoflurane and injected intravenously with 49 MBq ^{18}F -fluoride and 141 MBq $^{99\text{m}}\text{Tc}$ -MDP (not shown here). A 60 min whole-body acquisition was performed, starting 30 min post injection. Reconstructed images were post-filtered with a 0.8 mm FWHM 3D Gaussian.

2.6. Analysis

Time required for image reconstruction does not solely depend on the time per iteration of the algorithm, but also on the number of iterations needed to converge to the desired image. We quantify the convergence speed in the phantom experiment by calculating small detail contrast which is closely related to resolution. As the 0.8 mm rods are the smallest details still clearly visible in the resolution phantom for the collimator used, we use these to determine speed of convergence. The details of how contrast is calculated are explained below. To compare reconstruction times of different algorithms, we compare the times the different algorithms take to achieve equal contrast. As a measure of image quality, we perform a visual assessment of differently reconstructed images at equal contrast and we compare their noise levels. Details about noise level calculations are also provided below.

2.7. Hot-rod Derenzo phantom analysis

On (unfiltered) resolution phantom images reconstructed on 0.4 mm voxels, a contrast-noise analysis similar to the one presented in (Walker *et al* 2014) was done; the images were resampled to a fine grid and

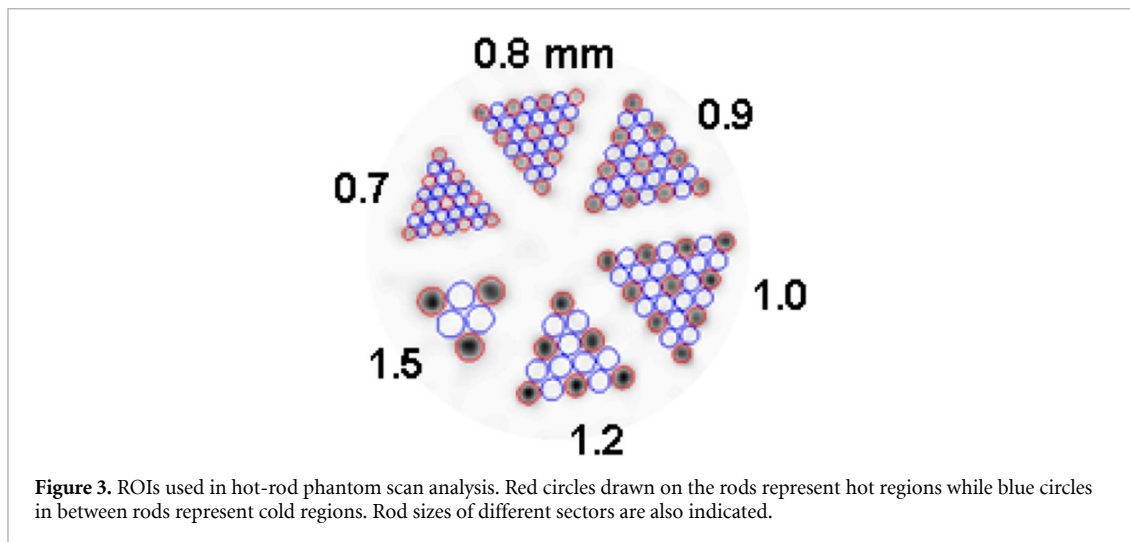


Figure 3. ROIs used in hot-rod phantom scan analysis. Red circles drawn on the rods represent hot regions while blue circles in between rods represent cold regions. Rod sizes of different sectors are also indicated.

regions-of-interest (ROIs) with a diameter of 0.9 times the rods sizes were then placed on and in-between the rods (see figure 3). ROI placement was repeated on subsequent 0.4 mm thick slices for a total number of 10 planes. For a certain sector, if \bar{h} is the mean activity in all ROIs placed on the rods and \bar{c} the mean activity in the ROIs placed in between (averaged over all slices), then contrast is defined to be

$$\text{Contrast} = \frac{\bar{h} - \bar{c}}{\bar{c}}.$$

Furthermore, to represent variability in ROI mean values we calculated for each sector the standard deviations σ_h and σ_c of the activity in ROIs on and in between rods, respectively. The noise in that sector is then calculated as

$$\text{Noise} = \sqrt{\sigma_h^2 + \sigma_c^2} / (\bar{h} + \bar{c})$$

2.8. Uniformly filled syringe analysis

As was found in (Goorden *et al* 2016), in case the tails were not modelled properly in reconstruction, part of the activity was reconstructed outside the syringe. To quantify if this is also the case for our proposed DM-DV approach, we calculated the activity in a cylindrical annulus with 19 mm inner diameter and 25 mm outer diameter around the 15.9 mm diameter phantom (in the same way as in the previous paper). Activity reconstructed in this region outside the phantom was reported as percentage of total reconstructed activity (for unfiltered images).

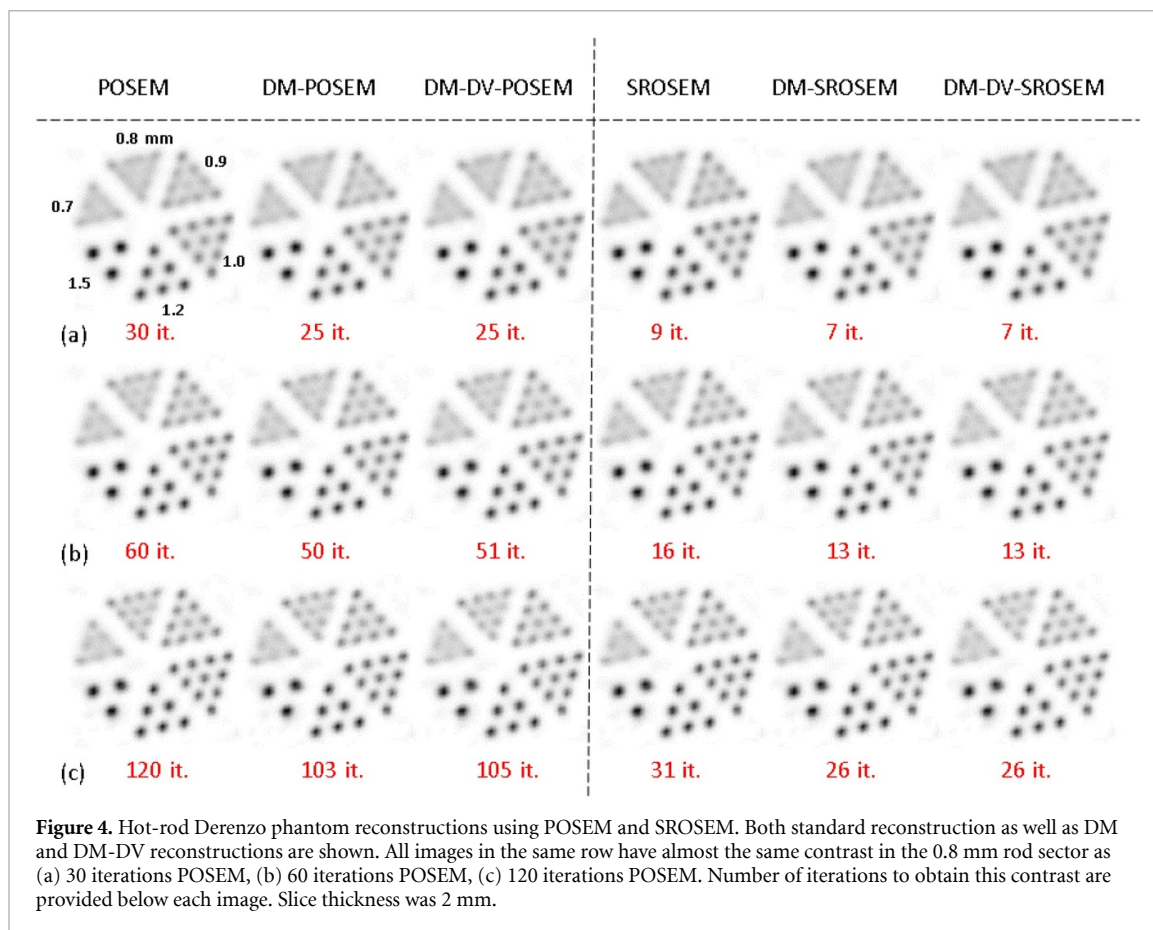
2.9. Timing

Reconstruction times provided in this paper were all obtained on our 24-core computer (Intel Xeon Gold 6126 Processors, 2.60 GHz, 19.25 M Cache). For the resolution phantom, average time per iteration is the average over the first 10 iterations. For all other scans, total reconstruction times were provided. We provide these reconstruction times both for 0.8 mm and 0.4 mm voxel size.

3. Results

In figure 4, images of the hot-rod Derenzo phantom are shown, reconstructed with POSEM or SROSEM, either combined with a DM approach (DM-POSEM and DM-SROSEM, respectively) or with a combined DM-DV approach (DM-DV-POSEM and DM-DV-SROSEM, respectively). We show three rows of reconstructed images, with the first image in each row being POSEM with 30 (figure 4(a)), 60 (figure 4(b)) and 120 iterations (figure 4(c)), respectively. The other images in the same row were selected to have equal contrast in the 0.8 mm rod sector as the contrast of the POSEM reconstruction. The number of iterations that were needed to obtain this contrast are shown underneath each image.

Figure 4 shows that the number of iterations required to converge to a certain contrast depends on the algorithm used. Here two effects play a role. First of all, convergence of reconstructions is somewhat faster when the DM approach is used instead of a standard reconstruction that uses the same matrix in forward



and backprojection. Such improved convergence properties for DM reconstruction were also reported in previous works, e.g. (Kamphuis *et al* 1998, Zeng and Gullberg 2000). Secondly, the use of SROSEM instead of POSEM decreases the number of required iterations further. The DV approach does not affect convergence speed. This is further quantified by the plots in figure 5 which show contrast vs. iteration number in the resolution phantom for all rod sectors. For the 0.8 mm rod sector, we performed a least squares fit to these curves to determine an equivalent iteration number for all tested algorithms. From this fit, we found that DM-POSEM and DM-DV-POSEM reach the same contrast in n iterations as POSEM does in $1.3 \cdot n$ iterations. Furthermore, contrast of SROSEM in n iterations is equal to that of $4.0 \cdot (n - 1)$ POSEM iterations while DM-SROSEM and DM-DV-SROSEM have an equivalent iteration number of $5.0 \cdot (n - 1)$. Note that the $(n - 1)$ term in the equivalent iteration number of the SROSEM algorithms is due to the fact that the first iteration in SROSEM is a single MLEM iteration which leads to negligible contrast recovery compared to POSEM with 32 subsets but which is necessary to determine the correct update scheme (Vaissier *et al* 2016). Furthermore, as the maximum number of subsets in SROSEM is 128, a speed up of SROSEM of 4 compared to POSEM is as expected at least in high-count regions where the maximum number of subsets is automatically used. Thus in case the number of iterations is $\gg 1$, SROSEM needs 4.0 times less iterations than POSEM while DM-DV-SROSEM requires 5.0 times less iterations. While these equivalent iteration numbers were obtained by fitting contrast curves for the 0.8 mm rod sector, figure 6 shows that they are very similar over different rod sectors; in this figure the same contrast curves as in figure 5 are plotted, but now against equivalent iteration number as defined by the formulas above. As a result, all curves fall on top of each other, indicating that convergence is accelerated similarly for different rod sizes.

Besides requiring less iterations to reach equal contrast, the DM-DV approach results in extra speed up as the time per iteration is reduced, both in the backprojection step (due to the DM approach) as in the forward projection step (because of the DV approach). The time required for one iteration is listed in table 1 for each of the algorithms. While in POSEM, a single iteration at 0.4 mm voxel size takes 617 s for the hot-rod phantom, with DM-DV-POSEM or DM-DV-SROSEM this time reduces to 161 s and 268 s, a 3.8 and 2.3 times reduction, respectively. The maximum acceleration factors over POSEM of all algorithms are provided in the final columns of table 1. These numbers are valid for the hot-rod phantom and for large numbers of iterations ($n \gg 1$) for which the first SROSEM iteration is negligible time-wise. From the table, it can be inferred that a large speed-up over POSEM can be attained; for 0.4 mm voxel size with DM-DV-POSEM, the

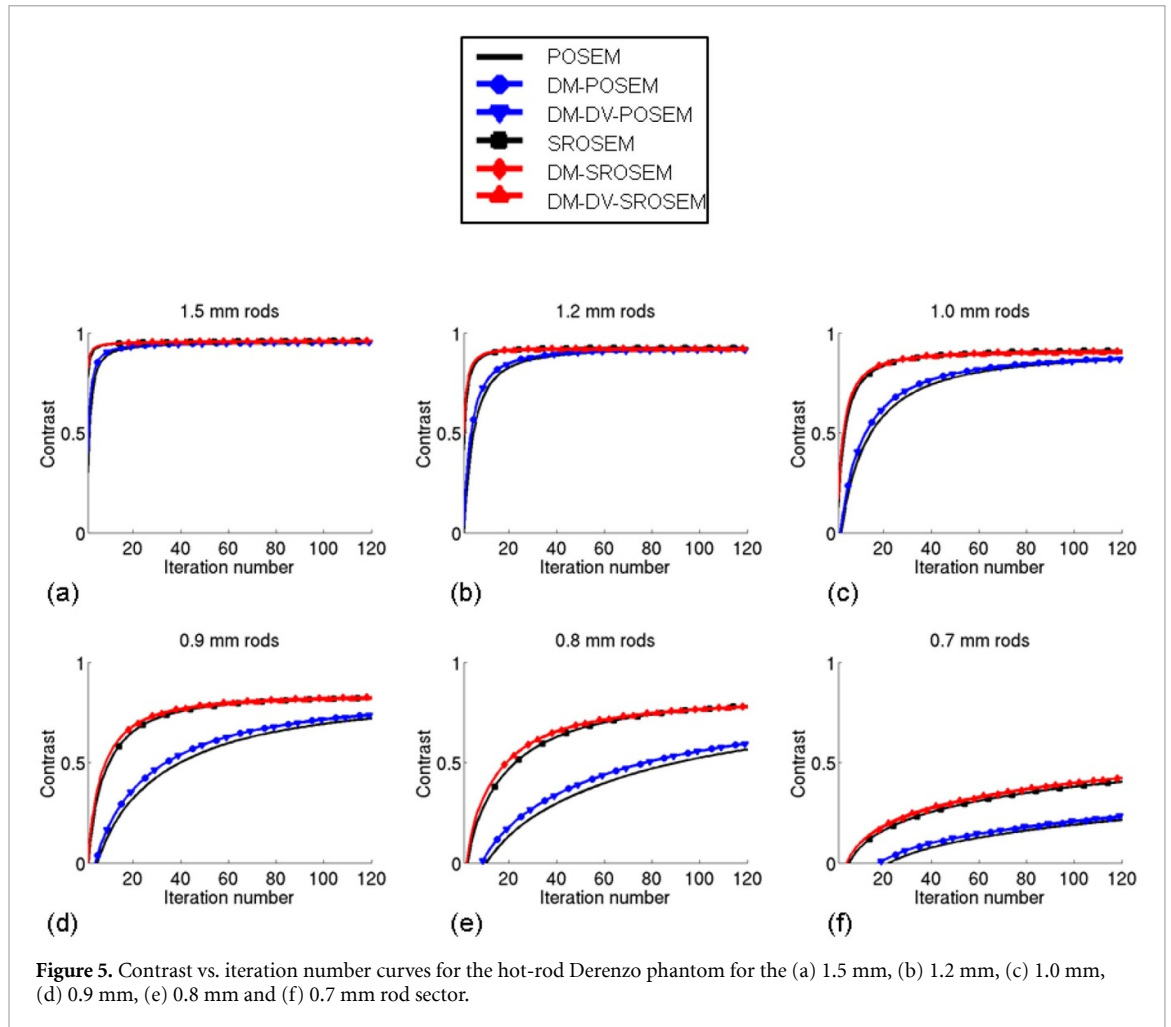


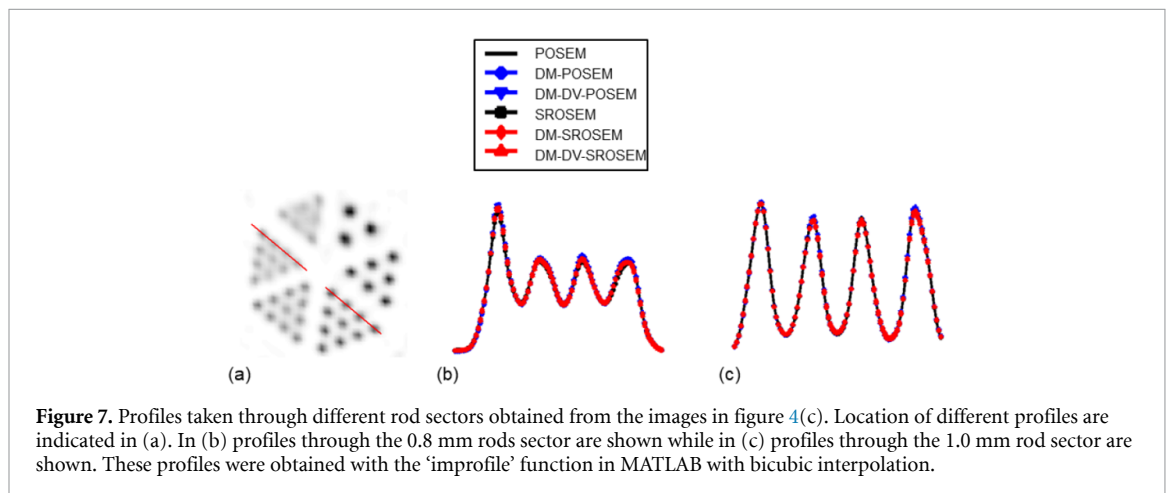
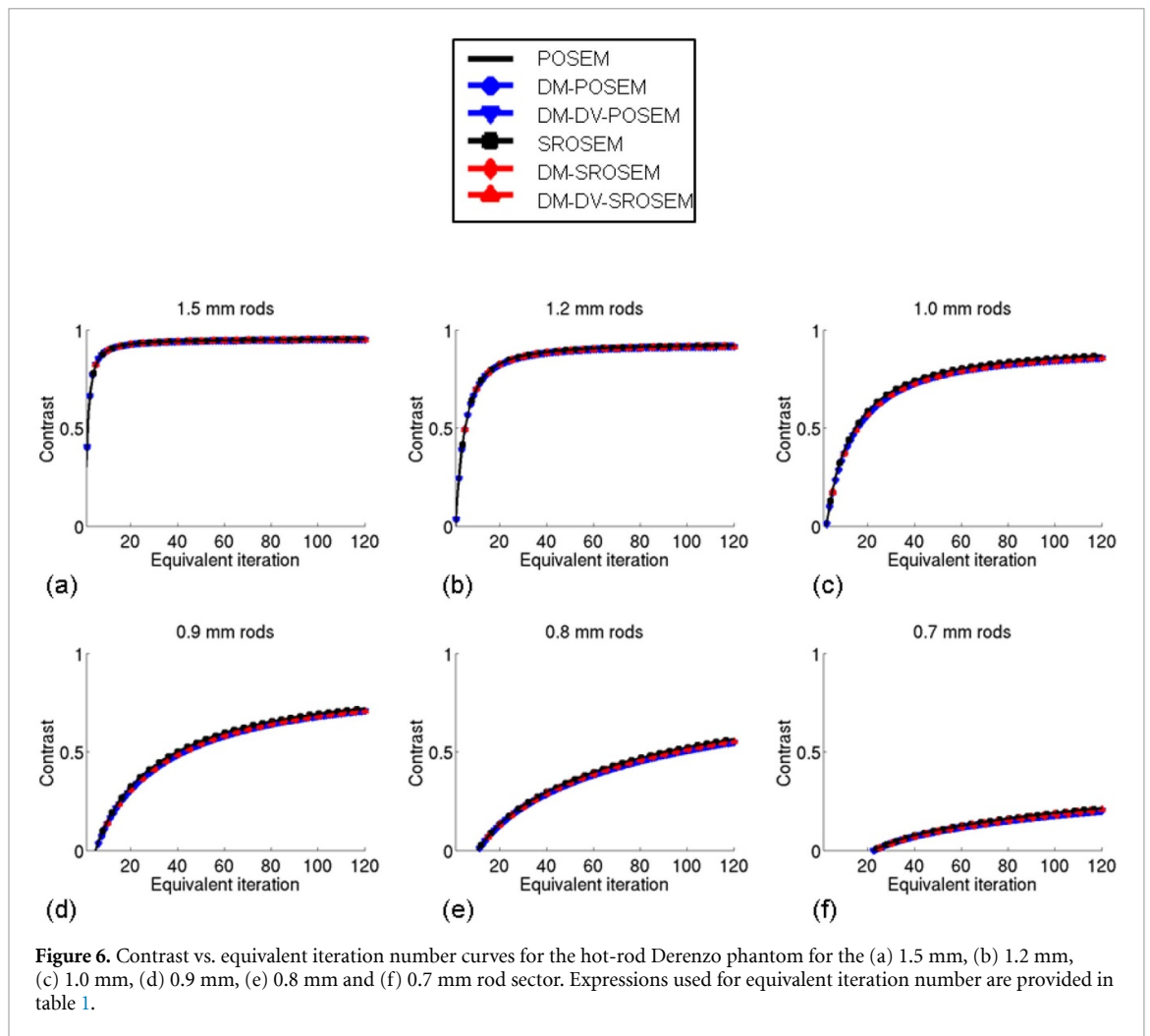
Figure 5. Contrast vs. iteration number curves for the hot-rod Derenzo phantom for the (a) 1.5 mm, (b) 1.2 mm, (c) 1.0 mm, (d) 0.9 mm, (e) 0.8 mm and (f) 0.7 mm rod sector.

Table 1. Overview of algorithms tested. For each algorithm, the equivalent iteration number compared to POSEM is provided (i.e. the number of POSEM iterations that would be required to reach the same contrast as reached after n iterations of the tested algorithm). Furthermore, the time per iteration (for the hot-rod phantom) is provided as well as the acceleration (in case $n \gg 1$), both for 0.4 mm and 0.8 mm voxel size.

Algorithm	Equiv. iteration number compared to POSEM	Time per iteration		Acceleration (for $n \gg 1$)	
		0.4 mm voxel	0.8 mm voxel	0.4 mm voxel	0.8 mm voxel
POSEM	n	617 s	80 s	1	1
DM-POSEM	$1.3 \cdot n$	286 s	37 s	2.8	2.8
DM-DV-POSEM	$1.3 \cdot n$	161 s	24 s	5.0	4.3
SROSEM	$4.0 \cdot (n - 1)$	703 s	97 s	3.5	3.3
DM-SROSEM	$5.0 \cdot (n - 1)$	380 s	56 s	8.1	7.1
DM-DV-SROSEM	$5.0 \cdot (n - 1)$	268 s	47 s	11.5	8.5

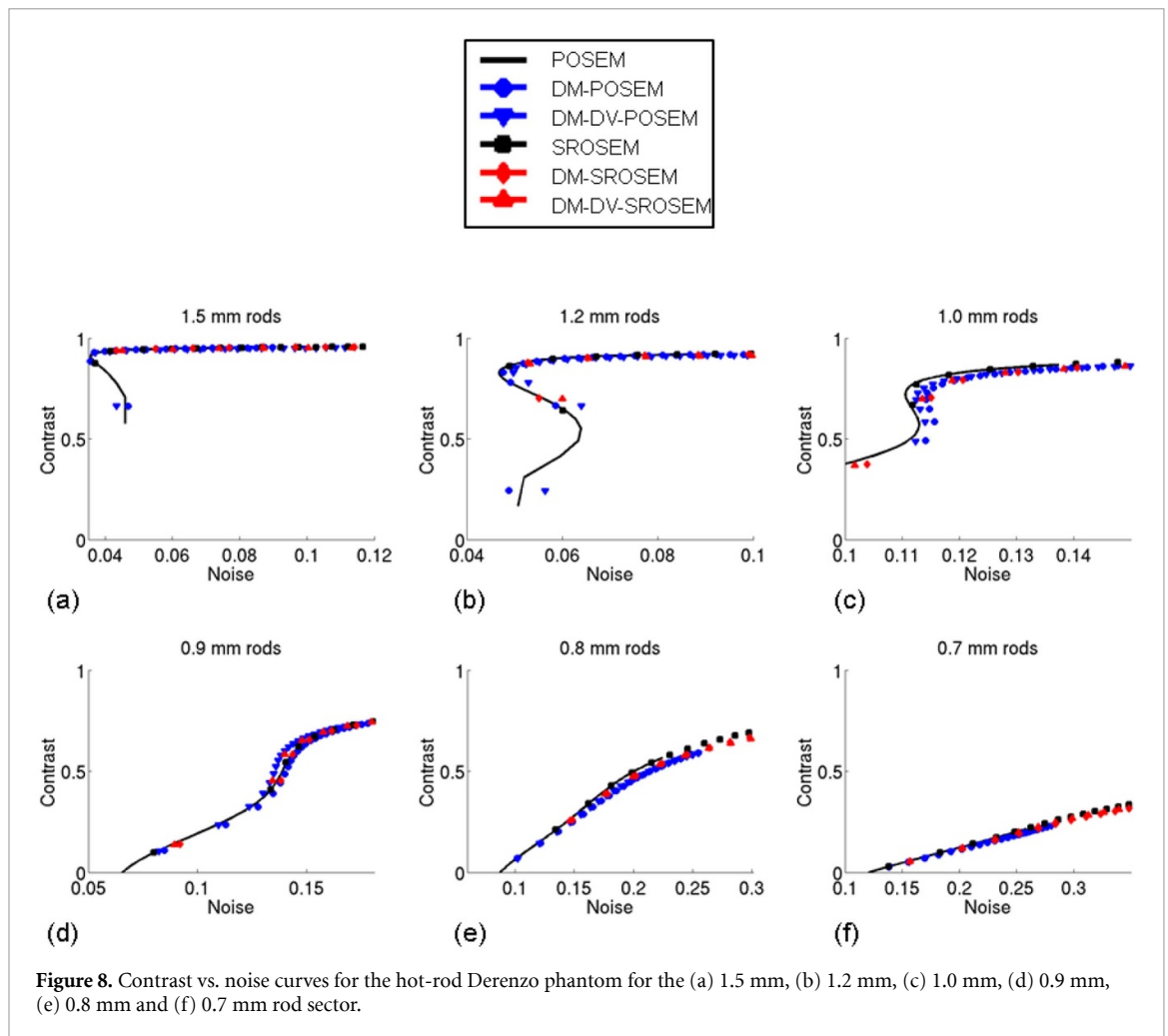
same contrast can be reached 5.0 times faster than with POSEM while with DM-DV-SROSEM the speed-up increases to 11.5. For the larger 0.8 mm voxel size, acceleration factors are somewhat reduced, probably because the overhead time is relatively more important for shorter reconstruction times.

After having established that the proposed algorithms accelerate image reconstruction, it is essential to test if reconstructed images with accelerated algorithms have the same quality as with standard POSEM. First of all, a visual assessment of reconstructed images in figure 4 does not reveal any visible difference for the hot-rod phantom images on the same row, meaning that images with equal contrast are visually indistinguishable. This is further confirmed by the profiles through the 0.8 mm and 1.0 mm rod sectors shown in figure 7. These profiles were taken from all images in figure 4(c) (i.e. for images with iteration number equivalent to 120 POSEM iterations). The figure shows that profiles are indistinguishable for different algorithms. To quantify similarity of images further, contrast as function of noise was plotted in figure 8, where the solid black line is the curve representing POSEM, while the symbols denote the other



tested algorithms. These figures show that images at equal contrast are very similar in terms of noise, at least once images are close to convergence (there are deviations in the earlier iterations).

Finally, figure 9 shows reconstructions of the uniformly filled syringe as well as several mouse studies reconstructed with POSEM and DM-DV-SROSEM (the fastest algorithm). Profiles through the images are provided as well for a more quantitative comparison. Images obtained with different algorithms have equivalent iteration number (determined by the equations in table 1). The total reconstruction time is provided below each image (both for 0.4 mm and 0.8 mm voxel size). These images as well as the image profiles reveal only small differences between different images; these small differences are sometimes visible between profiles and in the cardiac images the amount of activity in the apex is slightly different between OSEM and DM-DV-SROSEM. For the uniform phantom, we found that activity reconstructed outside the

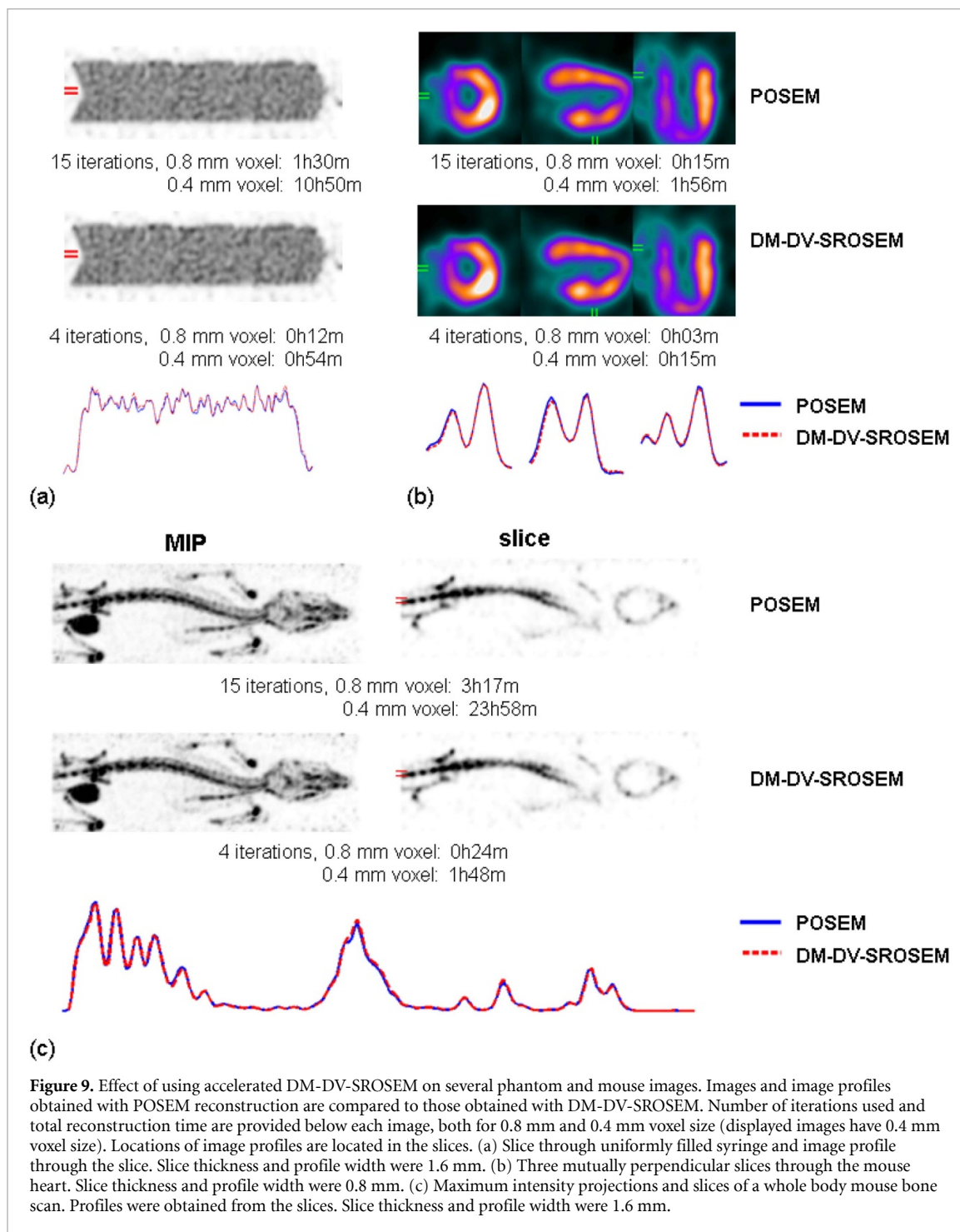


phantom was 4.8% for POSEM and 4.4% for DM-DV-SROSEM. In line with hot-rod phantom results, DM-DV-SROSEM achieves acceleration factors up till 13 over POSEM.

As accelerated algorithms all use a cut off value of 0.05 (to distinguish tails from central part of the PSF), one may wonder if simply using a cut off of 0.05 would provide sufficient image quality. In (Goorden *et al* 2016) we already investigated different cut offs and concluded that image quality was significantly improved by using a 0.01 cut off instead of 0.05 or higher. Therefore $C = 0.01$ was used as the gold standard here. This is also confirmed by the images we show in the [appendix](#) of this paper in which we compared results from this paper to POSEM and SROSEM reconstructed images with 0.05 cut off (denoted by POSEM(5%) and SROSEM(5%)). For the hot-rod phantom, using $C = 0.05$ leads to differences in profiles and negatively impacts contrast-noise characteristics as shown in figures A3 and A4 which are equivalent to figures 7 and 8 but with POSEM(5%) and SROSEM(5%) added. Visual differences are less clear in hot-rod images themselves (figure A2). Furthermore, in accordance with our previous work, using POSEM(5%) or SROSEM(5%) leads to increased background levels, especially for the uniform phantom and whole body mouse scan (figure A5). This is also confirmed quantitatively; for POSEM(5%) and SROSEM(5%), the respective amounts of activity reconstructed outside the phantom were 8.3% and 8.4%, approximately two times higher than the values reported for POSEM with a cut off value of 0.01 or for DM-DV-SROSEM.

4. Discussion

We presented a new approach for accelerated image reconstruction for VECTor-PET in which we combined an accelerated algorithm (pixel-based SROSEM) and a DM approach with a new DV method, which is particularly well-suited for long-tail PSFs as encountered in VECTor. With pixel-based DM-DV-SROSEM, we typically attained an order of magnitude shorter reconstruction times than with standard POSEM without any visible loss in image quality. Two factors play a role in this acceleration; first of all, both the use of SROSEM instead of POSEM as well as the use of a DM method accelerate image reconstruction as less iterations are required to reach the same level of contrast (by approximately a factor 5.0 for large numbers of



iterations). Secondly, the time per iteration is reduced; for the backprojection step this reduction is accomplished by using a smaller matrix with higher tail cut off, while for the forward projection step a rougher voxel size is used for those parts of the system matrix where probabilities are more slowly varying. Together this lead to an acceleration factor of 11.5 for the hot-rod phantom at 0.4 mm voxel size. For the animal images in some cases (e.g. the cardiac scan) somewhat lower acceleration factors were attained, partly due to the lower number of iterations required meaning that the first SROSEM iteration starts to play a larger role in total reconstruction time. Acceleration factors were also reduced for the smaller 0.8 mm voxel size, probably because in this case overhead time in the reconstruction software started to play a relatively larger role as the time spent on the projection and backprojection step quickly reduces as voxel size increases.

Comparison of images was done based on visual assessment, comparison of image profiles and a more quantitative contrast-noise analysis for the phantoms. Note that images were post-filtered with a Gaussian and image profiles were extracted from the filtered images. The amount of filtering was based on a visual assessment. Quantitative comparison was based on unfiltered images.

The DM approach that was applied here uses a backprojector with higher tail cut off than the forward projector, thus projector and backprojector in the reconstruction are unmatched. Such unmatched projector/backprojector pairs in which not all physics (e.g. scatter, attenuation, full system response) is included in the backprojector is usually applied to speed up reconstruction. In (Zeng and Gullberg 2000) a criterion for a valid pair was developed in terms of convergence, but authors also noted that for practical image reconstruction in which noise is present the use of a valid backprojector may not be critical. Here we take a heuristic approach in which we validate our DM approach by comparing several reconstructed images.

Reconstruction times reported in this paper were all obtained on the same 24-core computer that was available to us for an extended time (in order to perform all timings on the same work station). Of course, total reconstruction times could be further improved by using faster computers or computers with more cores. We already observed an almost twofold decrease in reconstruction time on other newer computers in our group, but could not perform our timings on these faster computers as they were heavily used by other researchers.

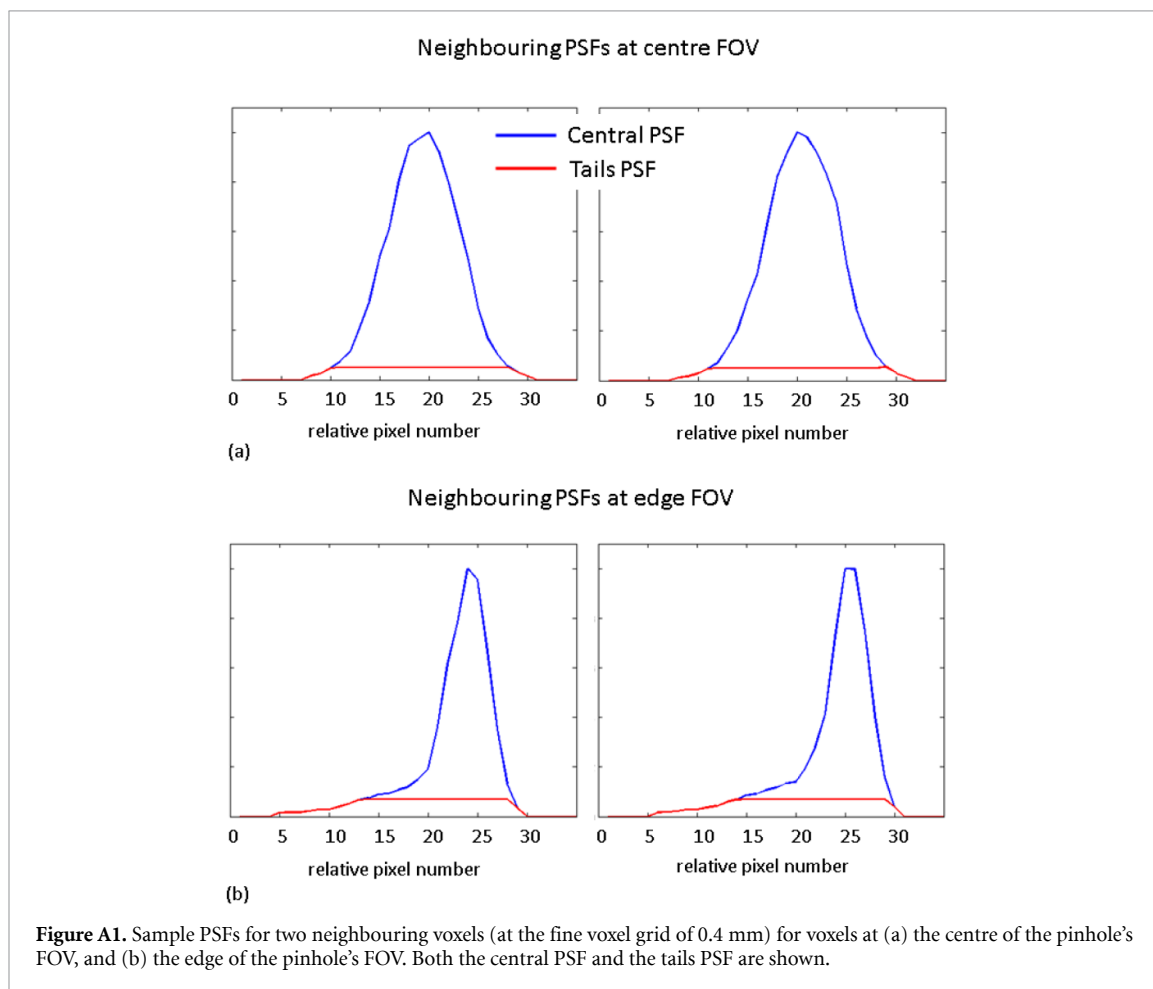
The amount of acceleration that is achieved depends on the choice of cut off parameters C_f and C_b . As mentioned, we chose these parameters based on preliminary try-outs, but did not perform any rigorous optimization. It may therefore be possible that further acceleration could be achieved if these parameters would further be optimized and higher values of these cut offs would be used. Another parameter that plays a role is the voxel size on which the tails are stored, which is at the moment three times rougher than the voxel size of the main part of the PSF. Note however that at the moment the 'tail matrix' has already more than five times fewer elements than the other matrices used in the DM-DV-SROSEM reconstructions (see section 2.3). Therefore, using an even rougher voxel size for the tails would probably not lead to much more acceleration.

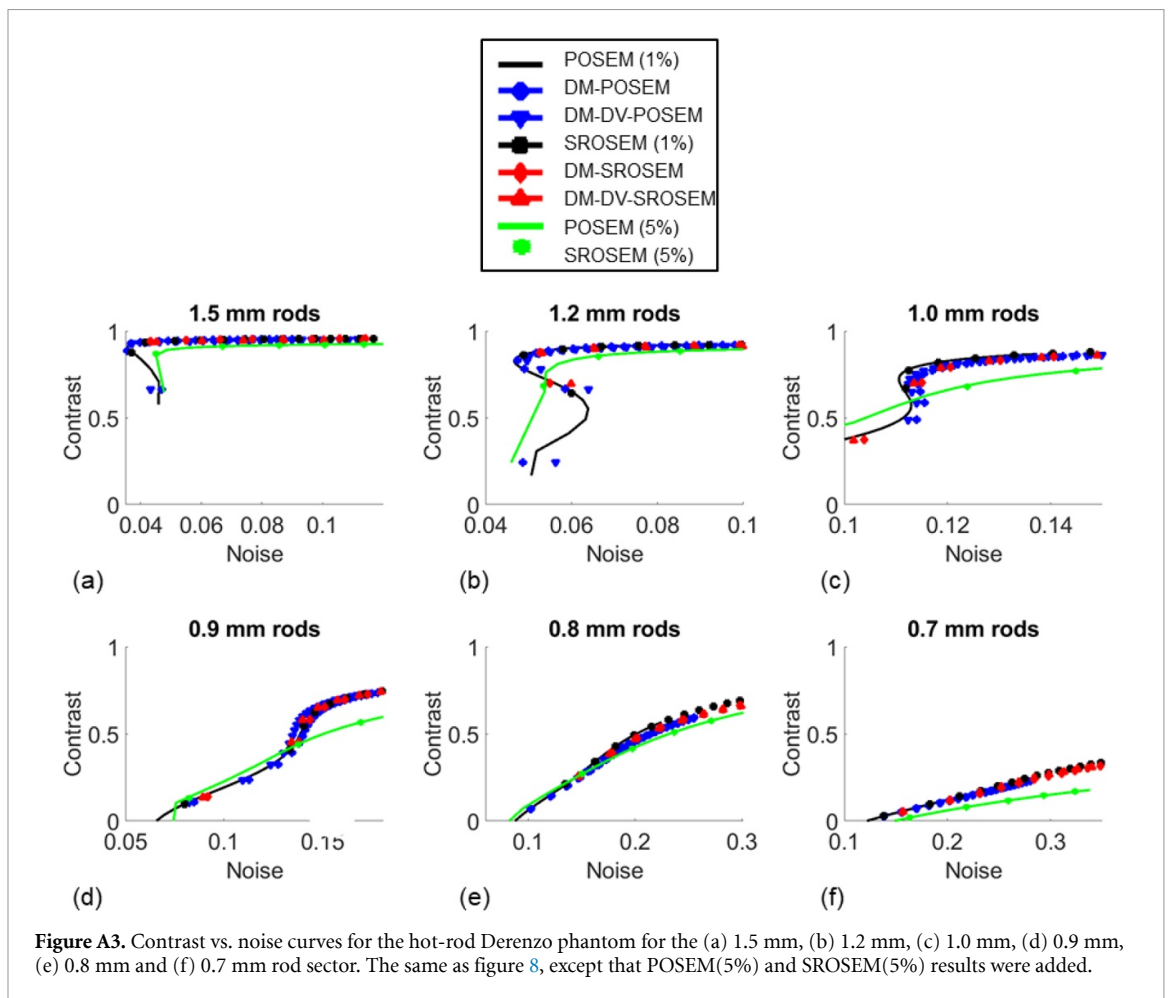
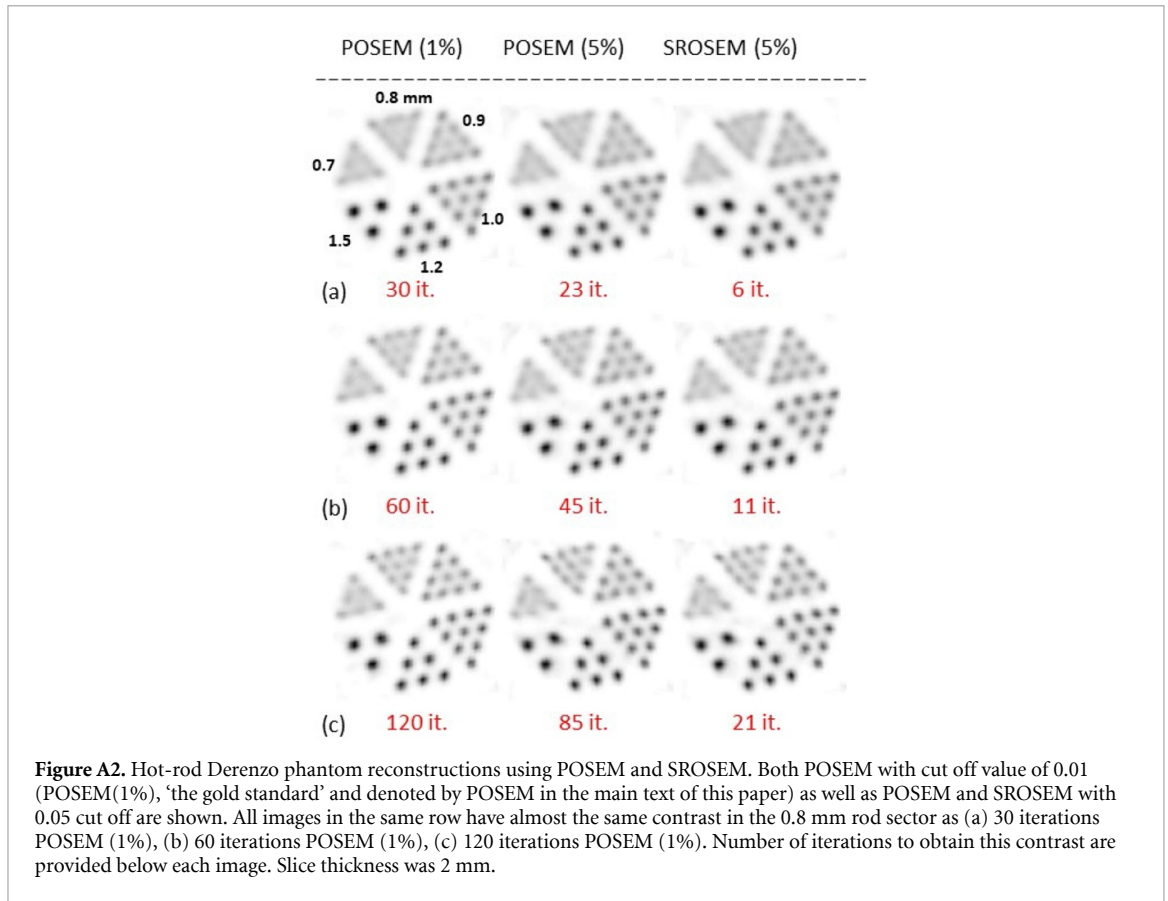
Although in this paper we focused on image reconstruction of PET tracer distributions (all studies used ^{18}F), we believe that this method is also very well suited to accelerate reconstruction for SPECT isotopes that emit high-energy gammas. It has already been shown that the clustered pinhole technique in VECTor is well-suited to image these isotopes, (e.g. de Swart *et al* 2016 for ^{213}Bi , van der Have *et al* 2016 for ^{131}I) and for their reconstruction long PSF tails need to be included as well. As the use of isotopes that also emit high-energy gammas is emerging (Herrmann *et al* 2017), mainly for theranostic applications, accurate and fast reconstruction methods can have many applications.

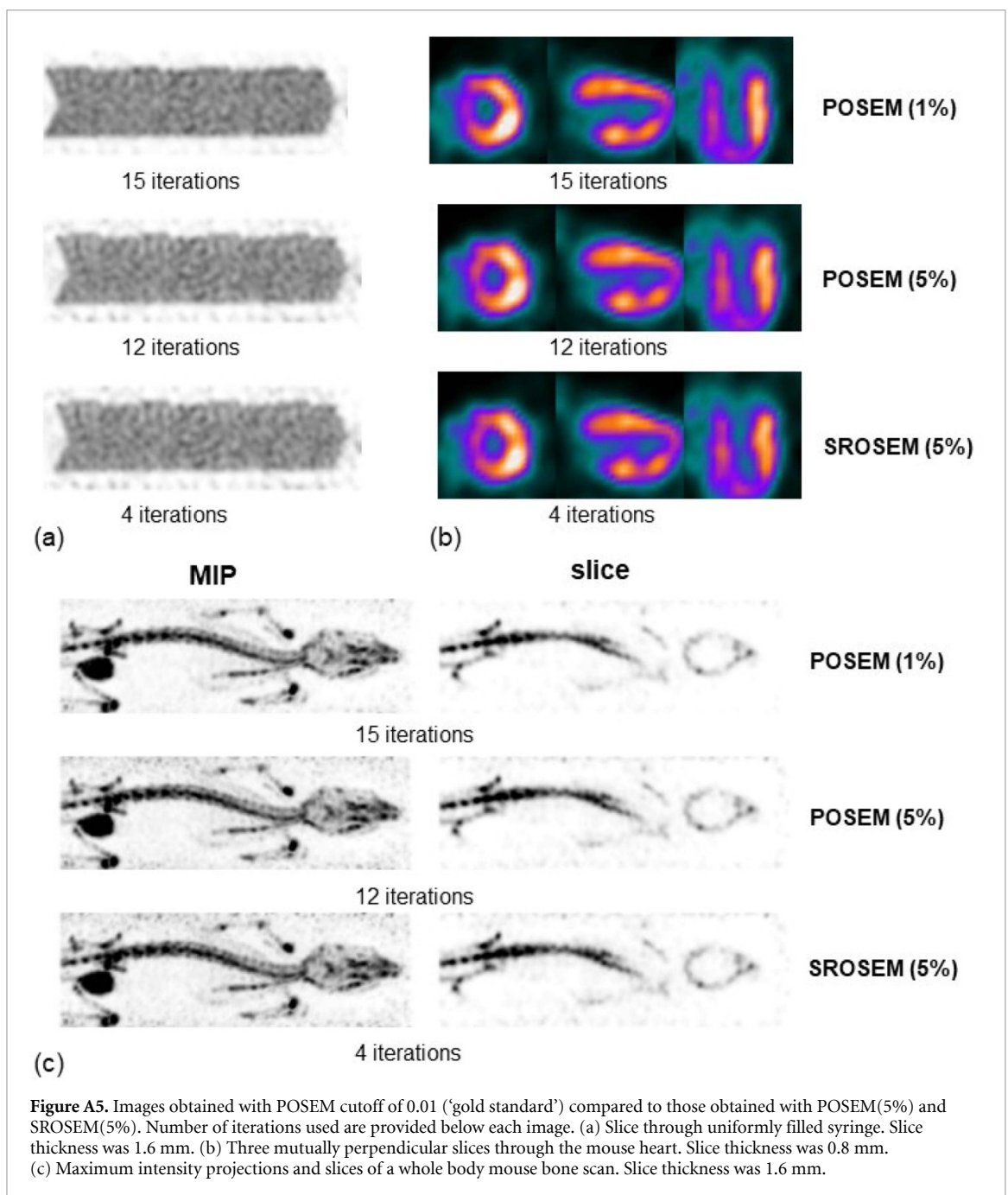
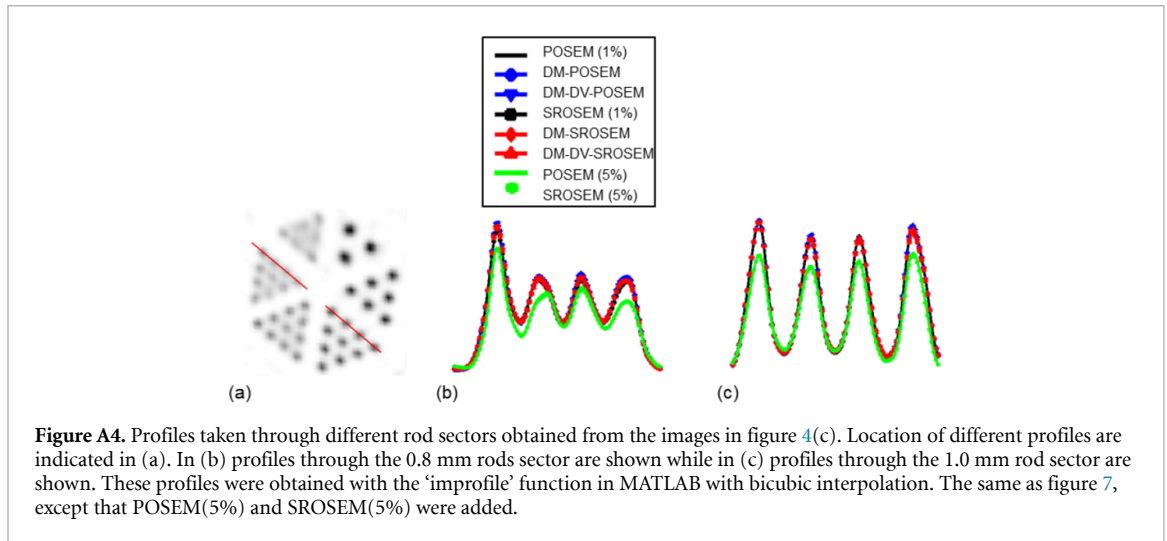
5. Conclusion

Compared to POSEM, DM-DV-SROSEM accelerates reconstruction by an order of magnitude while obtaining similar images. DM-DV-SROSEM thus enables accurate VECTor-PET image reconstruction by including photon transport in the PSF tails without inhibitive long reconstruction times.

Appendix







References

- Beekman F J 2011 Focused pinhole gamma detection device European Patent no. EP2073039, 2009 United States Patent US20090159802, 2011
- Beekman F J, de Jong H W A M and van Geloven S 2002 Efficient fully 3-D iterative SPECT reconstruction with Monte Carlo-based scatter compensation *IEEE Trans. Med. Imaging* **21** 867–77
- Beekman F J, Kamphuis C, Koustoulidou S, Vastenhouw B, Rendon D, Ramakers R M, Blazis S, van Cauter S, Goorden M C and Gremse F 2018 VECTor-6: broadband photon tomography takes the versatility of multi-modality preclinical imaging beyond the limits of PET-SPECT-CT and Optical Imaging *World Molecular Imaging Congress (WMIC) 2018 (Seattle, WA, 12–15 September 2018)*
- Branderhorst W, Vastenhouw B and Beekman F J 2010 Pixel-based subsets for rapid multi-pinhole SPECT reconstruction *Phys. Med. Biol.* **55** 2023
- Branderhorst W, Vastenhouw B, van der Have F, Blezer E L A, Bleeker W K and Beekman F J 2011 Targeted multi-pinhole SPECT *Eur. J. Nucl. Med.* **38** 552–61
- Chacko A M et al 2017 F-18-FDG as an inflammation biomarker for imaging dengue virus infection and treatment response *JCI. Insight* **2** e93474
- Chekol R et al 2018 (89)Zr-nimotuzumab for immunoPET imaging of epidermal growth factor receptor I *Oncotarget* **9** 17117–32
- Cherry S R et al 1997 MicroPET: A high resolution PET scanner for imaging small animals *IEEE Trans. Nucl. Sci.* **44** 1161–6
- Dalm S U et al 2017 Ga-68/Lu-177-NeobOMB1, a novel radiolabeled GRPR antagonist for theranostic use in oncology *J. Nucl. Med.* **58** 293–9
- de Swart J, Chan H S, Goorden M C, Morgenstern A, Bruchertseifer F, Beekman F J, de Jong M and Konijnenberg M W 2016 Utilizing high-energy gamma-photons for high-resolution Bi-213 SPECT in mice *J. Nucl. Med.* **57** 486–92
- Dogan M, Moysey S M J, Ramakers R M, Devol T A, Beekman F J, Groen H C and Powell B A 2017 High-resolution 4D preclinical single-photon emission computed tomography/x-ray computed tomography imaging of technetium transport within a heterogeneous porous media *Environ. Sci. Technol.* **51** 2864–70
- Esquinas P L, Rodriguez-Rodriguez C, De La Vega J C, Bokharaei M, Saatchi K, Shirmohammad M, Hafeli U O, Sossi V and Celler A 2017 ¹⁸⁸Re image performance assessment using small animal multi-pinhole SPECT/PET/CT system *Phys. Medica* **33** 26–37
- Goorden M C and Beekman F J 2010 High-resolution tomography of positron emitters with clustered pinhole SPECT *Phys. Med. Biol.* **55** 1265
- Goorden M C, van der Have F, Kreuger R, Ramakers R M, Vastenhouw B, Burbach J P H, Booij J, Molthoff C F M and Beekman F J 2013 VECTor: a preclinical imaging system for simultaneous submillimeter SPECT and PET *J. Nucl. Med.* **54** 306–12
- Goorden M C, van Roosmalen J, van der Have F and Beekman F J 2016 Optimizing modelling in iterative image reconstruction for preclinical pinhole PET *Phys. Med. Biol.* **61** 3712
- Herrmann K, Dahlbom M, Nathanson D, Wei L, Radu C, Chatziioannou A and Czernin J 2013 Evaluation of the Genisys4, a bench-top preclinical PET scanner *J. Nucl. Med.* **54** 1162–7
- Herrmann K, Larson S M and Weber W A 2017 Theranostic concepts: more than just a fashion trend-introduction and overview *J. Nucl. Med.* **58** 1s-2s
- Hudson H M and Larkin R S 1994 Accelerated image-reconstruction using ordered subsets of projection data *IEEE Trans. Med. Imaging* **13** 601–9
- Ivashchenko O, van der Have F, Villena J L, Groen H C, Ramakers R M, Weinans H H and Beekman F J 2015 Quarter-millimeter-resolution molecular mouse imaging with U-SPECT+ *Mol. Imaging* **14** 1–8
- Kadrmas D J, Frey E C, Karimi S S and Tsui B M W 1998 Fast implementations of reconstruction-based scatter compensation in fully 3D SPECT image reconstruction *Phys. Med. Biol.* **43** 857
- Kamphuis C, Beekman F J, van Rijk P P and Viergever M A 1998 Dual matrix ordered subsets reconstruction for accelerated 3D scatter compensation in single-photon emission tomography *Eur. J. Nucl. Med.* **25** 8–18
- Kim S et al 2019 Transneuronal propagation of pathologic α -synuclein from the gut to the brain models Parkinson's disease *Neuron* **103** 627–41
- King M A, Devries D J, Pan T S, Pretorius P H and Case J A 1997 An investigation of the filtering of TEW scatter estimates used to compensate for scatter with ordered subset reconstructions *IEEE Trans. Nucl. Sci.* **44** 1140–5
- Krishnamoorthy S, Blankemeyer E, Mollet P, Surti S, Van Holen R and Karp J S 2018 Performance evaluation of the MOLECUBES β -CUBE—a high spatial resolution and high sensitivity small animal PET scanner utilizing monolithic LYSO scintillation detectors *Phys. Med. Biol.* **63** 155013
- Lange R, de Klerk J M H, Bloemendal H J, Ramakers R M, Beekman F J, van der Westerlaken M M L, Hendrikse N H and Ter Heine R 2015 Drug composition matters: the influence of carrier concentration on the radiochemical purity, hydroxyapatite affinity and in-vivo bone accumulation of the therapeutic radiopharmaceutical ¹⁸⁸Rhenium-HEDP *Nucl. Med. Biol.* **42** 465–9
- Lecomte R, Cadorette J, Richard P, Rodrigue S and Rouleau D 1994 Design and engineering aspects of a high-resolution positron tomograph for small animal imaging *IEEE Trans. Nucl. Sci.* **41** 1446–52
- Miwa K, Inubushi M, Takeuchi Y, Katafuchi T, Koizumi M, Saga T and Sasaki M 2015 Performance characteristics of a novel clustered multi-pinhole technology for simultaneous high-resolution SPECT/PET *Ann. Nucl. Med.* **29** 460–6
- Adachi N et al 2017 In vivo simultaneous imaging of vascular pool and hypoxia with a HT-29 tumor model: the application of dual-isotope SPECT/PET/CT *Int. J. Sci.* **25** 26–39
- Nagy K, Toth M, Major P, Patay G, Egri G, Haggkvist J, Varrone A, Farde L, Halldin C and Gulyas B 2013 Performance evaluation of the small-animal nanoScan PET/MRI System *J. Nucl. Med.* **54** 1825–32
- Nguyen M P, Goorden M C, Kamphuis C and Beekman F J 2019 Evaluation of pinhole collimator materials for micron-resolution ex vivo SPECT *Phys. Med. Biol.* **64** 105017
- Qi J Y and Leahy R M 2006 Iterative reconstruction techniques in emission computed tomography *Phys. Med. Biol.* **51** R541
- Robertson A K H, Ramogida C F, Rodriguez-Rodriguez C, Blinder S, Kunz P, Sossi V and Schaffer P 2017 Multi-isotope SPECT imaging of the ²²⁵Ac decay chain: feasibility studies *Phys. Med. Biol.* **62** 4406
- Sanchez F et al 2012 Small animal PET scanner based on monolithic LYSO crystals: performance evaluation *Med. Phys.* **39** 643–53
- Sato K et al 2016 Performance evaluation of the small-animal PET scanner ClairvivoPET using NEMA NU 4-2008 Standards *Phys. Med. Biol.* **61** 696
- Surti S, Karp J S, Perkins A E, Cardi C A, Daube-Witherspoon M E, Kuhn A and Muehllehner G 2005 Imaging performance of A-PET: a small animal PET camera *IEEE Trans. Med. Imaging* **24** 844–52

- Szanda I, Mackewn J, Patay G, Major P, Sunassee K, Mullen G E, Nemeth G, Haemisch Y, Blower P J and Marsden P K 2011 National Electrical Manufacturers Association NU-4 performance evaluation of the PET component of the NanoPET/CT preclinical PET/CT scanner *J. Nucl. Med.* **52** 1741–7
- Ter Heine R *et al* 2014 Bench to bedside development of GMP grade Rhenium-188-HEDP, a radiopharmaceutical for targeted treatment of painful bone metastases *Int. J. Pharm.* **465** 317–24
- Vaissier P E B, Beekman F J and Goorden M C 2016 Similarity-regulation of OS-EM for accelerated SPECT reconstruction *Phys. Med. Biol.* **61** 4300
- Vaissier P E B, Goorden M C, Vastenhouw B, Van der Have F, Ramakers R M and Beekman F J 2012 Fast spiral SPECT with stationary γ -cameras and focusing pinholes *J. Nucl. Med.* **53** 1292–9
- van der Have F, Ivashchenko O, Goorden M C, Ramakers R M and Beekman F J 2016 High-resolution clustered pinhole ^{131}I iodine SPECT imaging in mice *Nucl. Med. Biol.* **43** 506–11
- van der Have F, Vastenhouw B, Rentmeester M and Beekman F J 2008 System calibration and statistical image reconstruction for ultra-high resolution stationary pinhole SPECT *IEEE Trans. Med. Imaging* **27** 960–71
- Vangestel C *et al* 2016 Preclinical evaluation of [^{111}In]MICA-401, an activity-based probe for SPECT imaging of *in vivo* uPA activity *Contrast Media Mol. Imaging* **11** 448–58
- Vastenhouw B and Beekman F 2007 Submillimeter total-body murine imaging with U-SPECT-I *J. Nucl. Med.* **48** 487–93
- Verhoog S, Kee C W, Wang Y, Khotavivattana T, Wilson T C, Kersemans V, Smart S, Tredwell M, Davis B G and Gouverneur V 2018 ^{18}F -Trifluoromethylation of unmodified peptides with 5- ^{18}F -(trifluoromethyl)dibenzothiophenium trifluoromethanesulfonate *J. Am. Chem. Soc.* **140** 1572–5
- Visser E P, Disselhorst J A, Brom M, Laverman P, Gotthardt M, Oyen W J G and Boerman O C 2009 Spatial resolution and sensitivity of the in vivo small-animal PET scanner *J. Nucl. Med.* **50** 139–47
- Walker M D, Goorden M C, Dinelle K, Ramakers R M, Blinder S, Shirmohammad M, van der Have F, Beekman F J and Sossi V 2014 Performance assessment of a preclinical PET scanner with pinhole collimation by comparison to a coincidence-based small-animal PET scanner *J. Nucl. Med.* **55** 1368–74
- Wang B, van Roosmalen J, Piet L, van Schie M A, Beekman F J and Goorden M G 2017 Voxelized ray-tracing simulation dedicated to multi-pinhole molecular breast tomosynthesis *Biomed. Phys. Eng. Express* **3** 045021
- Weissleder R, Ross B D, Rehemtulla A and Gambhir S S 2010 *Molecular Imaging: Principles and Practice* (Shelton, CT: People's Medical Publishing House–USA)
- Welch A and Gullberg G T 1997 Implementation of a model-based nonuniform scatter correction scheme for SPECT *IEEE Trans. Med. Imaging* **16** 717–26
- Yang Y F *et al* 2016 A prototype high-resolution small-animal PET scanner dedicated to mouse brain imaging *J. Nucl. Med.* **57** 1130–5
- Zeng G L and Gullberg G T 1992 Frequency-domain implementation of the 3-dimensional geometric point response correction in SPECT imaging *IEEE Trans. Nucl. Sci.* **39** 1444–53
- Zeng G S L and Gullberg G T 2000 Unmatched projector/backprojector pairs in an iterative reconstruction algorithm *IEEE Trans. Med. Imaging* **19** 548–55
- Zhang H, Bao Q A, Vu N T, Silverman R W, Taschereau R, Berry-Pusey B N, Douraghy A, Rannou F R, Stout D B and Chatziioannou A F 2011 Performance evaluation of PETbox: a low cost bench top preclinical PET scanner *Mol. Imaging Biol.* **13** 949–61
- Ziemons K *et al* 2005 The ClearPET (TM) project: development of a 2nd generation high-performance small animal PET scanner *Nucl. Instrum. Methods A* **537** 307–11