A multi-centre evaluation of eleven clinically feasible brain PET/MRI attenuation correction techniques using a large cohort of patients


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Available online 14 December 2016
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Aim: To accurately quantify the radioactivity concentration measured by PET, emission data need to be corrected for photon attenuation; however, the MRI signal cannot easily be converted into attenuation values, making attenuation correction (AC) in PET/MRI challenging. In order to further improve the current vendor-implemented MR-AC methods for absolute quantification, a number of prototype methods have been proposed in the literature. These can be categorized into three types: template/atlas-based, segmentation-based, and reconstruction-based. These proposed methods in general demonstrated improvements compared to vendor-implemented AC, and many studies report deviations in PET uptake after AC of only a few percent from a gold standard CT-AC. Using a unified quantitative evaluation with identical metrics, subject cohort, and common CT-based reference, the aims of this study were to evaluate a selection of novel methods proposed in the literature, and identify the ones suitable for clinical use.

Methods: In total, 11 AC methods were evaluated: two vendor-implemented (MR-AC_DIXON and MR-AC_UK), five based on template/atlas information (MR-ACSEG_BONE (Koesters et al., 2016), MR-ACONTARIO (Anazodo et al., 2014), MR-ACBOSTON (Izquierdo-Garcia et al., 2014), MR-ACUCL (Burgos et al., 2014), and MR-ACMAXPROB (Merida et al., 2015)), one based on simultaneous reconstruction of attenuation and emission (MR-AC_MLA (Benoit et al., 2015)), and three based on image-segmentation (MR-AC_MUNICH (Cabello et al., 2015), MR-ACCAR_RiDR (Juttukonda et al., 2015), and MR-ACRESOLUTE (Ladefoged et al., 2015)). We selected 359 subjects who were scanned using one of the following radiotracers: [18F]FDG (210), [11C]PiB (51), and [18F]florbetapir (98). The comparison to AC with a gold standard CT was performed both globally and regionally, with a special focus on robustness and outlier analysis.

Results: The average performance in PET tracer uptake was within ± 5% of CT for all of the proposed methods,
1. Introduction

Positron emission tomography/magnetic resonance imaging (PET/MRI) combines the powerful and functional imaging of PET for assessment of patients in oncology (Hillier et al., 2008) and neurology (Heiss, 2009) with MRI's excellent soft-tissue characterization, contrast, and power to provide additional functional information like perfusion and diffusion. A likely key clinical application of simultaneous PET/MRI is in neurological disorders, in particular in dementia (Bailey et al., 2015; Drzezga et al., 2014; Dukart et al., 2011). In order to accurately quantify the radioactivity concentration measured by PET, emission data need to be corrected for photon attenuation.

Currently, two fully integrated simultaneous whole-body human PET/MRI systems are commercially available: the Biograph mMR (Siemens Healthcare GmbH, Erlangen, Germany) and the Signa PET/MRI (GE Healthcare, Waukesha WI, USA). The vendor-implemented attenuation correction (AC) method in the Signa PET/MRI system is based on an atlas of MRI/CT pairs used to derive the approximate size and location of bones and air cavities (Wollenweber et al., 2013) but is yet to be fully evaluated on a large patient cohort (Sekine et al., 2016a). Two AC-methods are implemented in the Biograph mMR system. They are both segmenting the MR image into tissue classes: the two-point Dixon-Water-Fat sequence (Dixon) based on Martinez-Moller et al. (2009), which can lead to underestimation of PET tracer uptake near the skull (Andersen et al., 2014), and the ultrashort echo time sequence (UTE), based on a similar acquisition method as Catana et al. (2010) and a different segmentation method based on a voxel-based classifier, which is challenged by remaining segmentation errors in the skull base and near the frontal sinuses (Dickson et al., 2014). A third, atlas-based AC method is currently being developed (Koesters et al., 2016; Paulus et al., 2015).

To further improve the quantitative accuracy, a number of methods have been proposed in the literature. These can be categorized into three types: template/atlas-based, segmentation-based, and reconstruction-based. The template/atlas-based methods create a pseudo-CT image by co-registering database-subjects to a patient’s MR image. The atlas or template can be based on pairs of CT and dual UTE (Delso et al., 2014b; Roy et al., 2014), CT and anatomical MRI (Dixon, T1, and/or T2-weighted sequences) (Anazodo et al., 2014; Andreasen et al., 2015; Burgos et al., 2014; Izquierdo-Garcia et al., 2014; Merida et al., 2015; Schreibmann et al., 2010; Sekine et al., 2016a; Torrado-Carvajal et al., 2016). Some methods also use probabilistic measures or machine learning techniques to compare with an atlas or template (Chan et al., 2013; Chen et al., 2015; Hofmann et al., 2008; Johansson et al., 2011; Naivalpakkam et al., 2013; Poynton et al., 2014; Yang et al., 2013). Alternatives to template/atlas-based methods are the segmentation-based methods, which derive attenuation maps purely from MR images. These methods usually segment the T1 (Fei et al., 2012; Zaidi et al., 2003) or UTE images (Cabello et al., 2015; Catana et al., 2010; Juttukonda et al., 2015; Lade foged et al., 2015) into multiple tissue classes, each assigned a mean linear attenuation coefficient (LAC). Furthermore, a number of UTE-based methods calculate a patient specific bone density (Cabello et al., 2015; Juttukonda et al., 2015; Lade foged et al., 2015) using the normalized logarithmic difference between the UTE images (Ri map) (Keereman et al., 2010). A number of methods also exist employing special MRI sequences with modified sampling schemes using either short echo time (STE) or improved UTE sequences, using point-wise encoding time reduction with radial acquisition (PETRA) (Grodzki et al., 2012), using zero TE (ZTE) (Wiesinger et al., 2016), or using triple UTE (Aitken et al., 2014; Berkner et al., 2012; Su et al., 2015). Finally, maximum-likelihood reconstruction-based methods, where activity and attenuation are simultaneously reconstructed based on maximum-likelihood (Nuyts et al., 1999), using either time-of-flight information (TOF) (Mehranian and Zaidi, 2015; Rezaei et al., 2012, 2014; Salomon et al., 2011) or non-TOF information using MR-priors (Benoit et al., 2015), are yet to be evaluated for PET/MRI-brain applications. The methods using TOF would currently be limited to the Signa PET/MRI system or the sequential Ingenuity TF PET/MRI system (Philips Healthcare, Cleveland, OH) (Zaidi et al., 2011). However, recently, a TOF-based MLAA implementation failed to show additional benefit for attenuation correction in PET/MRI imaging compared to a robust atlas-based method (Mehranian et al., 2016).

The proposed methods demonstrate improvements compared to the currently vendor-implemented AC, and many studies report deviations in PET uptake after AC of only a few percent from a gold standard CT-AC; or conclude that the methods are ready for clinical use on the basis of quantitative accuracy (Burgos et al., 2014; Izquierdo-Garcia et al., 2014; Ladefoged et al., 2015). There are publications indicating the desire for more accuracy in PET brain attenuation correction (Andersen et al., 2014; Dickson et al., 2014; Hitz et al., 2014), but whether the problem of AC has been solved fully is still debatable (Bailey et al., 2015).

There is currently no consensus on a method for summarizing evaluations, and this lack of standardization of metrics makes direct

with the average ± SD global percentage bias in PET FDG uptake for each method being: MR-ACDIXON (−1.13 ± 3.5)%, MR-ACUTE (−5.7 ± 2.0)%), MR-ACCENTO (−4.3 ± 3.6)%, MR-ACMUNICH (3.7 ± 2.1)%, MR-ACMLAA (−1.9 ± 2.6)%, MR-ACSEGBONE (−1.7 ± 3.6)%, MR-ACUC (0.8 ± 1.2)%), MR-ACCAR-RdR (−0.4 ± 1.9)%, MR-ACMAXPROB (−0.4 ± 1.6)%, MR-ACBOSTON (−0.3 ± 1.8)%, and MR-ACRESOLUTE (0.3 ± 1.7)% ordered by average bias. The overall best performing methods (MR-ACBOSTON, MR-ACMAXPROB, MR-ACRESOLUTE and MR-ACUC, ordered alphabetically) showed regional average errors within ± 3% of PET with CT-AC in all regions of the brain with FDG, and the same four methods, as well as MR-ACCAR-RdR, showed that for 95% of the patients, 95% of brain voxels had an uptake that deviated by less than 15% from the reference. Comparable performance was obtained with PiB and florbetapir.

Conclusions: All of the proposed novel methods have an average global performance within likely acceptable limits (± 5% of CT-based reference), and the main difference among the methods was found in the robustness, outlier analysis, and clinical feasibility. Overall, the best performing methods were MR-ACBOSTON, MR-ACMAXPROB, MR-ACRESOLUTE and MR-ACUC, ordered alphabetically. These methods all minimized the number of outliers, standard deviation, and average global and local error. The methods MR-ACMUNICH and MR-ACCAR-RdR were both within acceptable quantitative limits, so these methods should be considered if processing time is a factor. The method MR-ACSEGBONE also demonstrates promising results, and performs well within the likely acceptable quantitative limits. For clinical routine scans where processing time can be a key factor, this vendor-provided solution currently outperforms most methods. With the performance of the methods presented here, it may be concluded that the challenge of improving the accuracy of MR-AC in adult brains with normal anatomy has been solved to a quantitatively acceptable degree, which is smaller than the quantification reproducibility in PET imaging.

C.N. Ladefoged et al.
NeuroImage 147 (2017) 346–359
comparisons across the published AC methods difficult. The methods rely on different gold/silver standards or make use of cohorts with different diseases or imaging tracers. The cohorts are of varying sizes, from 5 (Navalpakkam et al., 2013) to 92 subjects (Juttukonda et al., 2015), which might not be enough to ensure robustness and capture potential outliers in a clinical setting.

In this work, we present the performance of prototype methods within a multi-centre study, which include:

- A large patient cohort (n=359) who were scanned with three different radiotracers ([18F]fluoro-D-glucose (FDG), [11C] Pittsburgh-compound-B (PiB), and [18F]florbetapir), allowing for detailed evaluation of outliers and robustness.
- Patients recruited from clinical populations with established or emerging clinical indications for both PET and MRI, and therefore well-suited for evaluating regional effects of brain MR-AC.
- A unified comparison between 11 PET/MRI brain AC methods using identical metrics and a common CT-based reference: two vendor-implemented (Dixon and UTE), five based on template/atlas-based information (Anazodo et al., 2014; Burgos et al., 2014; Izquierdo-Garcia et al., 2014; Koesters et al., 2016; Merida et al., 2015), one-based on simultaneous reconstruction of attenuation and emission (Benoit et al., 2015), and three based on image-segmentation (Cabello et al., 2015; Juttukonda et al., 2015; Ladefoged et al., 2015).

The methods were assessed both globally and regionally using multiple metrics, including a detailed robustness and outlier analysis across the whole patient group. We also compared the methods by assessing the time that was required to execute the methods, referring to the actual run-time and scan-time of the corresponding sequences. While time may not be of high importance in a research setting, a clinical setup may demand high requirements for this factor in order to provide a timely diagnosis and acceptable patient management and throughput. The aim of this study was to evaluate the proposed methods and identify the ones suitable for clinical use by performing a unified quantitative evaluation.

2. Materials and methods

All patient studies were performed using a fully-integrated PET/MRI system (Siemens Biograph mMR, Siemens Healthcare, Erlangen, Germany) (Delso et al., 2011). For the purpose of obtaining a reference MRI system (Siemens Biograph mMR, Siemens Healthcare, Erlangen, Germany) (Delso et al., 2011). For the purpose of obtaining a reference MRI system (Siemens Biograph mMR, Siemens Healthcare, Erlangen, Germany) (Delso et al., 2011). For the purpose of obtaining a reference MRI system (Siemens Biograph mMR, Siemens Healthcare, Erlangen, Germany) (Delso et al., 2011).

2.2. Imaging protocol

2.2.1. PET

The PET scan information is given in Table 1. Patients were positioned head first with their arms down on the fully-integrated PET/MRI system. Data were acquired over a single bed position of 25.8 cm covering the head and neck for 10/20/15 min for FDG/PiB/ florbetapir respectively (Table 1 for initiation of scan). For the purpose of this study, the PET data from the PET/MR acquisition were reconstructed using 3D Ordinary Poisson-Ordered Subset Expectation Maximization (OP-OSEM) with 4/3/4 iterations, 21 subsets, and 3/5/5 mm Gaussian post filtering for FDG/PiB/florbetapir, respectively, on 344×344 matrices (2.1×2.1×2.0 mm³ voxels) in line with the respective clinical protocols.

2.2.2. MRI

The scan protocol included two-point Dixon images using the vendor-provided DIXON-VIBE AC sequence with repetition time (TR)/echo time 1 (TE1)/echo time 2 (TE2)=2300/1.23/2.46 ms, flip angle 10°, coronal orientation, voxel size of 2.6×2.6×3.12 mm³, 19 s acquisition time; a UTE AC sequence with TR/TE1/TE2=11.94/0.07/ 2.46 ms, a flip angle of 10°, axial orientation, field of view (FOV) of 300 mm², reconstructed on 192×192×192 matrices (1.6×1.6×1.6 mm³ voxels), 100 s acquisition time; and a T1w MPRAGE with TR/TE=1900/2.44 ms (FDG/PiB) and 2300/2.95 ms (florbetapir), inversion time=900 ms, flip angle=9°, sagittal orientation, reconstructed on 512×512×192 matrices (0.49×0.49×1 mm³ voxels) (FDG/PiB) and 256×240×176 matrices (1.05×1.05×1.2 mm³ voxels) (florbetapir), 300 s acquisition time. The patients injected with FDG or PiB were all imaged using the software version VB20P, whereas those injected with florbetapir were imaged with VB18P, which included a work-in-progress version of the UTE sequence. Due to the difference in version

<table>
<thead>
<tr>
<th>Tracer</th>
<th>Originating center</th>
<th>N (fat/water inverted)</th>
<th>Male/Female</th>
<th>Age (min-max) in years</th>
<th>Injected tracer (± SD) in MBq</th>
<th>Scan start p.i. (min-max) in min</th>
</tr>
</thead>
<tbody>
<tr>
<td>[18F]FDG</td>
<td>Rigshospitalet, Copenhagen Denmark</td>
<td>201 (9)</td>
<td>108/93</td>
<td>68 (23–96)</td>
<td>203 (± 20)</td>
<td>51 (24–134)</td>
</tr>
<tr>
<td>[11C]PiB</td>
<td>Rigshospitalet, Copenhagen Denmark</td>
<td>48 (1)</td>
<td>2424</td>
<td>68 (39–85)</td>
<td>426 (± 75)</td>
<td>43 (37–100)</td>
</tr>
<tr>
<td>[18F]florbetapir</td>
<td>Washington University, North Carolina, USA</td>
<td>88 (3)</td>
<td>21/67</td>
<td>71 (41–91)</td>
<td>353 (± 30)</td>
<td>0 n=75, or 52 (49–60) n=13</td>
</tr>
</tbody>
</table>

p.i.: post injection.
numbering, mainly expressed in the UTE sequences, the tracers were evaluated separately.

2.2.3. CT
A low dose CT (120 kVp, 36 mAs, 74 slices, 0.6×0.6×3 mm³ voxels) was acquired from each patient to be used as gold standard. The CT images were acquired either on the same day as the PET/MRI examination (FDG and PiB) or within 8.3 ± 6 days of each other with no surgical procedures in between (florbetapir).

2.3. Creating the attenuation maps
In all, nine proposed MR-AC methods were selected for evaluation. These were selected to be representative of the three major types of examination (FDG and PiB) or within 8.3 ± 6 days of each other with no surgical procedures in between (florbetapir).

Table 2
Jaccard distance for overlap with bone (μ > 0.101 cm⁻¹) segmented from MR-ACCT shown for each method and tracer. Results are shown for full head and further subdivided into top and bottom representing the area above and below the eyes. The best results are highlighted in bold. Note the Dixon method is left out due to there not being any bone present.

<table>
<thead>
<tr>
<th>Method</th>
<th>FDG</th>
<th></th>
<th></th>
<th>PiB</th>
<th></th>
<th></th>
<th>Florbetapir</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Full</td>
<td>Top</td>
<td>Bottom</td>
<td>Full</td>
<td>Top</td>
<td>Bottom</td>
<td>Full</td>
<td>Top</td>
<td>Bottom</td>
</tr>
<tr>
<td>UTE</td>
<td>0.56 ± 0.06</td>
<td>0.62 ± 0.10</td>
<td>0.51 ± 0.05</td>
<td>0.57 ± 0.06</td>
<td>0.65 ± 0.09</td>
<td>0.51 ± 0.04</td>
<td>0.32 ± 0.07</td>
<td>0.38 ± 0.12</td>
<td>0.27 ± 0.04</td>
</tr>
<tr>
<td>Segbone</td>
<td>0.54 ± 0.04</td>
<td>0.63 ± 0.07</td>
<td>0.48 ± 0.04</td>
<td>0.56 ± 0.03</td>
<td>0.65 ± 0.04</td>
<td>0.49 ± 0.03</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Ontario</td>
<td>0.60 ± 0.04</td>
<td>0.74 ± 0.05</td>
<td>0.52 ± 0.05</td>
<td>0.60 ± 0.05</td>
<td>0.73 ± 0.07</td>
<td>0.52 ± 0.04</td>
<td>0.63 ± 0.05</td>
<td>0.75 ± 0.07</td>
<td>0.55 ± 0.04</td>
</tr>
<tr>
<td>Boston</td>
<td>0.74 ± 0.06</td>
<td>0.79 ± 0.08</td>
<td>0.69 ± 0.07</td>
<td>0.78 ± 0.04</td>
<td>0.83 ± 0.05</td>
<td>0.73 ± 0.04</td>
<td>0.78 ± 0.03</td>
<td>0.81 ± 0.06</td>
<td>0.75 ± 0.03</td>
</tr>
<tr>
<td>UCL</td>
<td>0.80 ± 0.04</td>
<td>0.83 ± 0.05</td>
<td>0.77 ± 0.05</td>
<td>0.80 ± 0.04</td>
<td>0.83 ± 0.05</td>
<td>0.78 ± 0.04</td>
<td>0.82 ± 0.03</td>
<td>0.85 ± 0.04</td>
<td>0.80 ± 0.03</td>
</tr>
<tr>
<td>MaxProb</td>
<td>0.81 ± 0.04</td>
<td>0.85 ± 0.05</td>
<td>0.77 ± 0.04</td>
<td>0.82 ± 0.02</td>
<td>0.84 ± 0.04</td>
<td>0.79 ± 0.03</td>
<td>0.80 ± 0.03</td>
<td>0.83 ± 0.04</td>
<td>0.78 ± 0.03</td>
</tr>
<tr>
<td>MLA</td>
<td>0.59 ± 0.07</td>
<td>0.67 ± 0.09</td>
<td>0.53 ± 0.06</td>
<td>0.51 ± 0.07</td>
<td>0.60 ± 0.09</td>
<td>0.44 ± 0.06</td>
<td>0.36 ± 0.09</td>
<td>0.37 ± 0.15</td>
<td>0.35 ± 0.05</td>
</tr>
<tr>
<td>Munich</td>
<td>0.72 ± 0.04</td>
<td>0.85 ± 0.05</td>
<td>0.63 ± 0.05</td>
<td>0.73 ± 0.04</td>
<td>0.86 ± 0.05</td>
<td>0.64 ± 0.04</td>
<td>0.72 ± 0.03</td>
<td>0.85 ± 0.04</td>
<td>0.63 ± 0.04</td>
</tr>
<tr>
<td>CAR-RiDR</td>
<td>0.50 ± 0.07</td>
<td>0.62 ± 0.09</td>
<td>0.40 ± 0.06</td>
<td>0.50 ± 0.06</td>
<td>0.62 ± 0.08</td>
<td>0.41 ± 0.06</td>
<td>0.46 ± 0.06</td>
<td>0.60 ± 0.09</td>
<td>0.36 ± 0.05</td>
</tr>
<tr>
<td>RESOLUTE</td>
<td>0.68 ± 0.04</td>
<td>0.85 ± 0.04</td>
<td>0.56 ± 0.04</td>
<td>0.69 ± 0.03</td>
<td>0.84 ± 0.04</td>
<td>0.58 ± 0.04</td>
<td>0.68 ± 0.03</td>
<td>0.85 ± 0.03</td>
<td>0.54 ± 0.04</td>
</tr>
</tbody>
</table>

Fig. 1. Attenuation correction images for a sample patient that minimizes the difference of the overall brain error to the median error across all methods. (A) CT, (B) Dixon, (C) UTE, (D) Segbone, (E) Ontario, (F) Boston, (G) UCL, (H) MaxProb, (I) MLA, (J) Munich, (K) CAR-RiDR, (L) RESOLUTE.
MR-AC categories (template/atlas-based, segmentation-based, and reconstruction-based), and had to be applicable to the Siemens mMR data available. Furthermore, they should be implemented on a central site (Copenhagen). Twelve attenuation maps were created. First, vendor-provided MR-based attenuation maps (MR-ACDIXON and MR-ACUTE) were derived using the DIXON VIBE sequence and the UTE MR sequence. Next, the nine proposed brain MR-AC methods were used. A brief introduction of each method is given below. We refer to the original publications for detailed explanations. Finally, for each subject, the CT image was co-registered to the T1 image using a 6-parameter rigid alignment procedure (minctracc, McConnell Imaging Center, Montreal, Canada) with normalized mutual information as objective function. The patient bed and head-holder were extracted manually from the CT images using an oval ROI (OsiriX software) applied to each transverse CT slice. The attenuation values were then converted from HUs to LACs at 511 keV by using a standard bi-linear scaling approach (Carney et al., 2006) as implemented in the Siemens PET/CT systems used. The co-registered CT attenuation map was substituted into the UTE attenuation file (MR-ACUTE) to facilitate use of routine reconstruction. We blurred the resulting image with a 4 mm Gaussian filter to simulate PET/CT reconstructions. Due to the limited coverage in the neck region by the acquired CT, we replaced the missing area by the values from MR-ACDIXON. To ensure a fair comparison, this replacement was also performed in all the other attenuation maps.

All attenuation maps, except for MR-ACDIXON, were created in the

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All attenuation maps, except for MR-ACDIXON, were created in the
image space defined by the method, and post-resampled and placed into the UTE file just as the reference CT. Since the CT image was co-registered to the T1 image, we also co-registered the Dixon and UTE images to the T1 image, and aligned the MR-AC maps of the methods defined using either of these.

All of the methods presented in this study, except for two, had been made available and were processed in Copenhagen. The MR-ACSEGBOONE attenuation maps were created by Siemens, and MR-ACUCU attenuation maps were created in London, as the CT/MRI database could not be shared at the time of this study.

2.3.1. Template/Atlas-based methods (Segbone, Ontario, Boston, UCL, MaxProb)

- **Siemens Healthcare GmbH - Segbone**: The prototype method by Siemens Healthcare (Koesters et al., 2016; Paulus et al., 2015) is a template-based method, where an MR model image of the skull region is non-rigidly registered to an individual subject exclusively on the Dixon input images. A pre-aligned skull mask is then brought to subject space, and attenuation values greater than soft-tissue are superimposed on the original Dixon attenuation map. This method is still a work-in-progress. The attenuation map is denoted MR-ACSEGBOONE.

- **London Ontario**: The method by Anazodo et al. (2014) uses the SPM8 function ‘new segment’ to extract a bone probability map from each individual’s T1 MPRAGE image. The bone map defined by voxels with probability above 80% are, following a post-processing procedure, superimposed on MR-ACDIXON and assigned the constant value 0.143 cm⁻¹. The resulting attenuation map is denoted MR-ACONTARIO.

- **Boston MGH**: The method by Burgos et al. (2014) non-linearly aligns a database of 41 T1-CT pairs to the intensity-normalized T1 image of a patient. At each voxel, a patch is extracted and the local normalized correlation coefficient is calculated between the patient T1 and each of the T1 images in the database. A weighting vector is calculated from the coefficients ranking each T1-CT pair relative to the other pairs, and used to obtain a target CT voxel value. The resulting synthesized CT image is converted to LACs, blurred with a 4 mm Gaussian filter, and denoted MR-ACBOSTON.

- **London UCL**: The method by Burgos et al. (2014) non-linearly aligns a database of 41 T1-CT pairs to the intensity-normalized T1 image of a patient. At each voxel, a patch is extracted and the local normalized correlation coefficient is calculated between the patient T1 and each of the T1 images in the database. A weighting vector is calculated from the coefficients ranking each T1-CT pair relative to the other pairs, and used to obtain a target CT voxel value. The resulting synthesized CT image is converted to LACs, blurred with a 4 mm Gaussian filter, and denoted MR-ACUCL.

- **Lyon - MaxProb**: The method by Merida et al. (2015) is similar to Burgos et al. (2014) by also aligning a database of 27 T1-CT pairs to an intensity-normalized T1 MPRAGE image of a patient, but deviates in the method used to combine the database into a target CT image. The authors discretize the CT images into air, soft tissue and bone voxels, calculate the modal tissue type across the database, and assign the mean of the CT values from the database subjects with that tissue type to the voxel. The final fused image is converted to LACs, blurred with a 4 mm Gaussian filter, and denoted MR-ACMAXPROB.

2.3.2. Maximum-likelihood reconstruction-based

- **Copenhagen - MLAA**: The method by Benoit et al. (2015) is a reconstruction-based method aimed at optimizing the existing non-TOF MLAA technique. The method uses the individual’s MR-ACUTE and a discretized version of the individual’s T1 image as prior information to the MLTR algorithm. A heuristically determined schema assigns the value for αj based on the two prior images, as well as the update term calculated in each MLTR iteration step. The final image of the MLAA algorithm is denoted MR-ACMLAA.

2.3.3. Segmentation-based (Munich, CAR-RiDR, RESOLUTE)

- **Munich**: The method by Caballo et al. (2015) uses the individual’s UTE TE1 and TE2 images to extract air, soft tissue as well as the bone signal, which is extracted using R2* and scaled to LACs by normalizing to the maximum intensity and thresholding using an empirically chosen value. The intensity of the bone voxels from the R2* map are equalized by the mean and maximum values from a CT-based database. The resulting image is blurred with a 3 mm Gaussian filter, and denoted MR-ACMUNCH.

- **University of North Carolina - CAR-RiDR**: The method by Juttukonda et al. (2015) also uses the individual’s UTE TE images to extract air, soft tissue and R2* signal. By fitting a sigmoid-best-fit to a set of 98 R2*-CT relationship pairs, the authors convert the measured R2* signal to a continuous CT value. The resulting image is denoted MR-ACCAR-RI DR.

- **Copenhagen - RESOLUTE**: The method by Ladefoged et al. (2015) segments brain, CSF, soft tissue, and air from the UTE TE images, and also uses R2* to extract a bone signal, which is converted to LACs using a set of R2*-CT relationship pairs from 10 training patients. To limit possible bias from regions with known R2* noise (Delso et al., 2014a), and because the amount of included bone is higher compared to Caballo et al. (2015), Juttukonda et al. (2015), the threshold for included bone was regionally varied within the head. The resulting image is blurred with a 4 mm Gaussian filter and denoted MR-ACRESOLUTE.

2.4. Image analysis

As Dixon-based methods may suffer from fat/water inversion (Ladefoged et al., 2014), we excluded the patients with inversion from further analysis in most of the metrics. This was the case for nine FDG, one PiB, and three florbetapir patients. To avoid bias in the results, the patients with fat/water inversion are excluded in most figures. When included, it will be clearly stated. The methods directly affected by the inversion are MR-ACSEGBOONE, MR-ACONTARIO and MR-ACCAR-RIDR.

2.4.1. Global attenuation map performance

To evaluate the performance precision of the methods based on the attenuation maps, we measured the Jaccard-index for the bone as being greater than 0.101 cm⁻¹, representing values above soft-tissue.

2.4.2. Global PET performance

As a prior step to evaluating the global PET performance, we first created individual brain masks. We moved all patient data to common MNI space using ANTs (Avants et al., 2011) by non-rigidly registering the patients’ T1 images to the ICBM 152 2009a template (Fonov et al., 2009). The MNI brain mask was back-warped to patient space, and the PET image reconstructed with the CT attenuation map was thresholded at > 20% of the maximum intensity value. Finally, the intersection between the two masks was calculated and used as the final mask of the brain, and used throughout this manuscript when referring to the brain region. Next, the voxel-wise percent difference relative to PETCT, defined as:

\[
\text{Rel}_C = \frac{\text{PET}_C - \text{PET}_{CT}}{\text{PET}_{CT}} \times 100,
\]

as well as the absolute percent difference, defined as:

\[
\text{Abs}_C = \frac{|\text{PET}_C - \text{PET}_{T1}|}{\text{PET}_{CT}} \times 100,
\]

were calculated for the PET images corrected with each of the evaluated
We calculated the averaged performance of $\text{Rel}_\%$ and $\text{Abs}_\%$ for the full brain across all patients for each of the methods. We computed the distribution of $\text{Rel}_\%$ errors for each of the methods using a histogram analysis. Finally, we computed the voxel-wise correlation between $\text{PET}_{CT}$ and each of the proposed methods for all voxels within the brain. We plotted the correlations in a joint histogram, and

![joint_histograms](image)

**Fig. 4.** Summed joint histograms of PET activity within the brain mask for FDG (n=192, excluding patients with fat/water tissue inversion) for $\text{PET}_{CT}$ versus each of the methods. The $R^2$ scores are average ± SD of the individual patients. The joint entropy (JE) is calculated for all patients.
computed the goodness-of-fit value ($R^2$) to the identity line for each of the patients.

2.4.3. Regional PET performance

To evaluate the regional PET performance, we back-warped the anatomical predefined regions from MNI space (Collins et al., 1999; Fonov et al., 2009) to each patient, and extracted the mean $Rel_{\text{BG}}$ for each region in the template. We furthermore calculated the average and standard deviation $Rel_{\text{BG}}$-images across all patients for each of the methods for visual evaluation.

2.4.4. Analysis of robustness and outliers

To estimate the number of outliers for each method, we introduced a novel metric. The metric calculates the percentage of patients with at least $x\%$ of the brain within $\pm y\%$ in the $Rel_{\text{BG}}$ images, where $x$ varies from 0–100 and $y\,=\,(5,10,15)$. A perfect method therefore has 100% of the patients within $\pm 0\%$ in the $Rel_{\text{BG}}$ images. To understand the reason for the worst outliers, we then analyzed the three FDG patients with the worst performance based on the $\pm 10\%$ threshold and categorized the error. We present the worst outlier of each method.

3. Results

3.1. Global attenuation map evaluation

Fig. 1 shows the axial, sagittal and coronal views for each attenuation method for a single sample patient. The representative sample patient was chosen objectively to be that which minimizes the difference to the median $Rel_{\text{BG}}$ error (Eq. (1)) across all methods.

The Jaccard similarity coefficient for bone for each tracer-group was comparable. For the largest group (FDG) the averaged scores ranged between 0.50–0.81 for the entire head, 0.40–0.77 in the bottom of the head covering from the eye socket and downwards, and 0.51–0.85 for the top of the head. The overall best performance was seen for MR-ACBOSTON, MR-ACMAXPROB, MR-ACMUNICH, MR-ACRESOLUTE and MR-ACUCL, in alphabetical order, with Jaccard indices of 0.68–0.81 in the entire head, and 0.79–0.85 in the top of the head. The individual result for all tracer-groups and methods are shown in Table 2. Note that MR-ACSEGBOBone was not evaluated on florbetapir-data, and thus no results are available.

3.2. Global PET performance

The reconstructed PET images for the sample patient in Fig. 1 are shown for the same slice in Fig. 2. Visually comparing the maximum uptake areas in PETCT, it is apparent that PETDIXON and PETUTE underestimate the PET uptake, whereas all the alternative methods improve the performance. For this patient, the average error for the entire brain was between $−3.3\%$ (Ontario) and 3.9% (Munich) for the proposed methods, and $−10\%$ and $−5.3\%$ for Dixon and UTE, respectively. The global $Rel_{\text{BG}}$ performance for the entire brain is shown for all 201 FDG patients and 11 methods in the boxplot in Fig. 3. The average performance was within $\pm 5\%$ of PETCT for all of the proposed methods, with the best performance seen for MR-ACBOSTON, MR-ACCAR-RiDR, MR-ACMAXPROB, MR-ACRESOLUTE, and MR-ACUCL, ordered alphabetically. MR-ACDIXON showed $−11\%$ and MR-ACUTE showed $−7\%$. The global $Abs_{\text{BG}}$ performance for the entire brain is summarized for each method and tracer in Supplementary Table 1. The global histogram analysis is shown in Supplemental Figs. 1–3 for the three tracers. The averaged voxel-wise joint histograms are shown for PETCT versus each of the proposed methods in Fig. 4 (FDG) and Supplemental Figs. 4–5 (PiB and Florbetapir). The highest correlation with the unity line was for FDG obtained by MR-ACRESOLUTE (0.96 ± 0.02), MR-ACUCL (0.96 ± 0.02), MR-ACMAXPROB (0.96 ± 0.04), MR-ACBOSTON (0.95 ± 0.03), and MR-ACCAR-RiDR (0.94 ± 0.03).

3.3. Regional PET performance

The regional $Rel_{\text{BG}}$ performance for the anatomical regions defined in the MNI template is shown in Fig. 5 for FDG and Supplemental Figs. 6–7 for PiB and Florbetapir. The best performance across all regions was again found with MR-ACBOSTON, MR-ACCAR-RiDR, MR-ACMAXPROB, MR-ACRESOLUTE and MR-ACUCL, ordered alphabetically. Only MR-ACMAXPROB and MR-ACRESOLUTE, ordered alphabetically, had an average $± 1\%$ error across all tracers below $± 5\%$ in all regions. The averaged absolute difference, $Abs_{\text{BG}}$, is shown in Supplemental Figs. 8–9 for each of the regions and tracers. The average and standard deviation images of the $Rel_{\text{BG}}$ calculated for each FDG patient are shown in Figs. 6 and 7, respectively, and in Supplemental Figs. 10–13.
for PiB and florbetapir. Considering all regions in the brain and all tracers, the overall best performance in terms of error and robustness was seen for MR-ACBOSTON, MR-ACMAXPROB, MR-ACRESOLUTE and MR-ACUCL, ordered alphabetically.

3.4. Analysis of robustness and outliers

The result of the outlier analysis is shown in Fig. 8 for errors within ± 5%, ± 10%, and ± 15% (FDG) and Supplemental Fig. 14 (PiB and florbetapir). The proposed methods exceed the performance of the current Dixon and UTE methods. The cutoff where all of the patients are within ± 10% of PET CT is achieved for MR-ACBOSTON, MR-ACMAXPROB, MR-ACRESOLUTE and MR-ACUCL ordered alphabetically, and at 90% of the brain, and between 60% and 80% for the other methods. At ± 15%, there is a clear separation between five of the methods (MR-ACBOSTON, MR-ACCAR-RiDR, MR-ACMAXPROB, MR-ACRESOLUTE and MR-ACUCL ordered alphabetically) and the remaining methods. The robustness of each method can be assessed in the standard deviation images (Fig. 7). The best performance was obtained by MR-ACUCL, in the center of the brain, shortly followed by MR-ACMAXPROB, MR-ACBOSTON and MR-ACRESOLUTE. In the cortical regions, the best performance was obtained by MR-ACRESOLUTE and MR-ACCAR-RiDR. The three patients with the worst performance are categorized in Table 3, and the worst outlier is shown for each method in Supplemental Fig. 15.

In general, the worst outliers in the template/atlas-based methods (MR-ACONTARIO, MR-ACBOSTON, MR-ACUCL, MR-ACMAXPROB) all have the same error category – the bone density is over or underestimated. Only a single case is due to registration errors (#1 in MR-ACMAXPROB). The effect of the error is greatest in the cortical regions, with errors up to 30% locally.

The general errors in the proposed segmentation-based methods (MR-ACMUNICH, MR-ACCAR-RiDR, MR-ACRESOLUTE) are over/under estimation of bone in air/tissue interface areas, resulting in local errors of up to 25% in the medulla and cerebellum. For the vendor-provided methods, the worst outliers were due to missing bone (MR-ACDIXON) and underestimated bone density (MR-ACUTE).

4. Discussion

This study facilitates a framework for comparing a large number of state-of-the-art methods using multiple metrics, and does so across a large patient cohort. We aimed to represent a broad spectrum of published methods, showing promises of clinical feasibility, and
selected nine methods that were applicable to our large cohort of Siemens mMR examinations. Adding the vendor-provided methods, Dixon and UTE, we evaluated 11 methods for obtaining an MR-AC image. The key findings for the proposed methods were: almost all achieved relative errors below ± 5% globally and in all regions of the brain; the methods essentially differed in robustness, clinical feasibility, and outliers. Dixon and UTE have been included as current vendor references and confirm earlier results from Andersen et al. (2014), Dickson et al. (2014) with a tendency to global underestimation of activity.

The cerebellum is a region that is often used for normalization purposes, especially in kinetic modeling, and therefore requires extra attention. Considering the average ± 1 SD, the best performance is obtained with MR-ACBOSTON and MR-ACMAXPROB, shortly followed by most of the other methods; the only methods exceeding ± 5% were MR-ACMLAA and MR-ACMUNICH (Fig. 5). The maximum error in the cerebellum was 6–9% from the reference for most of the proposed methods (Supplemental Fig. 16), except for MR-ACCAR-RiDR (14%), MR-ACMLAA and MR-ACMUNICH (17%).

The result of the attenuation map evaluation (Table 2) illustrates that the multi-atlas and multi-registration based methods (MR-ACUCL, MR-ACMAXPROB) have the highest overall bone accuracy, with equally good results achieved in the upper part of the head by R2*-based methods (MR-ACMUNICH, MR-ACRESOLUTE). This area-dependent difference could be explained by the challenges reported in air/tissue interface areas originating from the UTE sequence, such as the mouth and esophagus (Delso et al., 2014a). Metal implant-induced artifacts resulting in overestimation of bone in the mouth area especially affects the accuracy of the R2*-based methods, but obviously also the remaining methods. Of all methods, two of the R2*-based methods (MR-ACCAR-RiDR and MR-ACRESOLUTE) had the lowest standard deviation in the cortical regions near the bone (Fig. 7). This can be explained by the fact that these methods extract patient specific bone density, whereas the template/atlas-based methods use an average CT value. This can in some cases lead to local under- or overestimation of the PET signal of up to 20% (Table 3, Supplemental Fig. 15). It should be noted that for the subjects where the CT include the dental region, the CTs were also affected by metal implants resulting in streak artifacts, which could bias the results further.

The template/atlas-based methods generally produced attenuation maps very similar to the CT-based attenuation maps, but they are based on the assumption that any new subject can be represented by a template, a local combination of atlas patients, or by a non-rigid registration to a template. The template/atlas-based methods are
therefore also potentially challenged by abnormal anatomy and patient surgery. The template/atlas-based methods are, furthermore, currently only optimized and tested for adults, as the application to children might require a purpose specific MRI/CT database. We did not include children, or any patients who have undergone anatomy-changing surgery in this study.

Most subjects chosen came from dementia studies, as this disease has the potential to be a key clinical application of PET/MRI (Bailey et al., 2015). The disease is clinically imaged with FDG, or amyloid binding radiotracers such as PiB or florbetapir, with the majority of patients having relatively widely distributed uptake in the brain, making the tracers well suited for a large-scale evaluation of MR-AC methods. The overall performance was, generally, similar across the three tracers for all methods (Supplemental Figs. 1–3).

When considering patient management, the acquisition time and number of steps required for generating AC maps has also to be taken into account. A potential challenge for the template/atlas-based methods is the long computation time due to multiple registrations for each patient in the database (Table 4). The MR-ACMAXPROB and MR-ACUCL methods currently require at least an hour of processing time for each attenuation map, whereas the MR-ACMUNICH and MR-ACCAR-RiDR methods only take a few seconds. The shorter processing time is usually a key argument for choosing the segmentation-based methods in fast-paced clinical services, whereas accuracy is much more important than speed in research applications. The MR-ACSEGbone prototype requires less than 2 min of processing time, with further potential for speed-up. The processing time for multi-atlas methods could be shortened by code optimization and by reducing the number of database studies using pre-selection of the candidates more likely to match the patient (e.g. using gender, age, etc.) (Aljabar et al., 2009) but this requires further evaluation. An alternative is to use the newly proposed one-registration multi-atlas approach, which, however, comes at the cost of a lower accuracy in bone extraction compared to the multi-atlas approach (Arabi and Zaidi, 2016). In line with this, recent result on GE Signa system has shown, that compared to the clinical single-atlas, multi-atlas (MR-ACUCL) is more accurate especially in regions close to the skull base (Sekine et al., 2016b). The emission-based method, MLAA, is currently computationally intensive, and relies on multiple MR sequences, which might complicate its use in a clinical setting.

We excluded a single machine-learning-based AC-method (Chen et al., 2015) from this study, as it had excessively long processing time for each patient (>$4$ days), making it infeasible both in clinical routine and in this study in its current state.

4.1. Limitations

The use of a CT as gold standard is debatable. The CT image does not fully cover the FOV of the MR and PET area, and is further subject to streak artifacts from dental implants. Since the CT images are not simultaneously acquired with the PET/MRI, they are also subject to non-rigid movement, e.g. jaw movement and swallowing, not accounted for during the rigid registration. In lack of a true gold standard, such as a transmission scan with a rotating source, we considered the CT to be the reference.

The cohort, mainly consisting of patients with dementia and age-matched controls, represents an ageing population. This study did not include data to test the performance of these MR-AC methods on brain scans from younger patients. Furthermore, the study cohort did not include patients with focal PET lesions. Therefore, the performance of these methods in brain tumor patients has yet to be evaluated in most cases, although evaluation of individual methods (e.g. MR-ACBOSTON) has already been performed in challenging brain tumors as in Izquierdo-Garcia et al. (2014).

The data from Washington University (florbetapir) were used to train the mapping of bone values from HU to LACs in the MR-ACCAR-RiDR method (Juttukonda et al., 2015). This could potentially bias the results.

5. Conclusion

All of the proposed methods have an average global performance within likely acceptable limits ($\pm 5\%$ of CT-based reference), and
differences between methods were found in terms of the robustness, clinical feasibility and outlier analysis. The overall best performance was seen in the MR-ACBOSTON, MR-ACMAXPROB, MR-ACRESOLUTE and MR-ACUCL methods, ordered alphabetically, which all minimize the number of outliers, standard deviation, and average global and local error. The methods MR-ACCAR-RiDR and MR-ACMUNICH were both within acceptable quantitative limits, so these methods should be considered if processing time is a factor. The method MR-ACSEGBONE also demonstrates promising results, and performs well within the likely acceptable quantitative limits. For a clinical routine setting where time is critical, this vendor-provided prototype solution currently outperforms most methods, especially when acquisition time of the fast DIXON is included. With the performance of the methods presented here, it may be concluded that the challenge of improving the accuracy of MR-AC in adult brains with normal anatomy has been solved to an acceptable degree, with errors smaller than the quantification reproducibility in PET imaging.

Acknowledgements

A special thanks to biomedical engineering student David Nyrnberg and computer science student Casper Hansen for aiding with the large data processing tasks involved in this study. The PET/MR system at Rigshospitalet was kindly provided by the John and Birthe Meyer Foundation, Denmark. At UCL this work was supported by an IMPACT studentship funded jointly by Siemens and the UCL Faculty of

Table 3
Analysis of worst three outlier patients based on amount of brain with errors greater than ± 10% (excluding patients with fat/water inversion). The worst outlier is also shown in Supplemental Fig. 15.

<table>
<thead>
<tr>
<th>Subject with highest % of brain &gt; ±10%</th>
<th>2nd highest % of brain &gt; ±10%</th>
<th>3rd highest % of brain &gt; ±10%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dixon</td>
<td>83% (A)</td>
<td>82% (A)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>−30% (cortical)</td>
<td>−20% (cortical)</td>
</tr>
<tr>
<td>UTE</td>
<td>58% (A)</td>
<td>55% (A)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>−15% (cortical)</td>
<td>−15% (cortical)</td>
</tr>
<tr>
<td>Segbone</td>
<td>35% (A)</td>
<td>28% (B)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>−15% (cortical)</td>
<td>+20% (cortical)</td>
</tr>
<tr>
<td>Ontario</td>
<td>44% (A)</td>
<td>39% (A)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>−20% (cortical)</td>
<td>−20% (cortical)</td>
</tr>
<tr>
<td>Boston</td>
<td>10% (B)</td>
<td>10% (B)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>20% (cortical)</td>
<td>10% (cortical)</td>
</tr>
<tr>
<td>UCL</td>
<td>10% (B)</td>
<td>9% (B)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>30% (cortical)</td>
<td>15% (cortical)</td>
</tr>
<tr>
<td>MaxProb</td>
<td>14% (D)</td>
<td>12% (A)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>−25% (r brain)</td>
<td>−20% (cortical)</td>
</tr>
<tr>
<td>MLAA</td>
<td>34% (A)</td>
<td>34% (A)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>−15% (cortical)</td>
<td>−15% (cortical)</td>
</tr>
<tr>
<td>Munich</td>
<td>41% (C)</td>
<td>34% (C)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>25% (cerebellum)</td>
<td>20% (cerebellum)</td>
</tr>
<tr>
<td>CAR-RiDR</td>
<td>23% (A)</td>
<td>23% (C)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>−20% (cortical)</td>
<td>20% (cerebellum)</td>
</tr>
<tr>
<td>RESOLUTE</td>
<td>10% (B/C)</td>
<td>9% (B/C)</td>
</tr>
<tr>
<td>Rel_% (region)</td>
<td>10% (cerebellum)</td>
<td>10% (cerebellum)</td>
</tr>
</tbody>
</table>

Category explanation: (A) Bone density or bone width underestimated, (B) bone density or bone width overestimated, (C) incorrect air/bone interface segmentation, and (D) template/atlas registration errors.

Table 4
Requirements of the proposed methods.

<table>
<thead>
<tr>
<th>T1 MPRAGE (acq. time: 300 s)</th>
<th>Dixon (acq. time: 19 s)</th>
<th>UTE (acq. time: 100 s)</th>
<th>Prior data</th>
<th>Requirements (license)</th>
<th>Runtime (approx.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Segbone</td>
<td>✓</td>
<td>✓</td>
<td>Needs single atlas</td>
<td>Proprietary</td>
<td>1–2 m</td>
</tr>
<tr>
<td>Ontario</td>
<td>✓</td>
<td>✓</td>
<td>Needs single atlas</td>
<td>FreeSurfer, SPM (GNU)</td>
<td>30 m</td>
</tr>
<tr>
<td>Boston</td>
<td>✓</td>
<td>✓</td>
<td>Needs multiple atlases</td>
<td>NiftyReg (BSD)</td>
<td>1 h</td>
</tr>
<tr>
<td>UCL</td>
<td>✓</td>
<td>✓</td>
<td>Needs multiple atlases</td>
<td>NiftyReg (BSD)</td>
<td>2 h</td>
</tr>
<tr>
<td>MaxProb</td>
<td>✓</td>
<td>✓</td>
<td>Optimization needed per tracer type</td>
<td>Required</td>
<td>1 h</td>
</tr>
<tr>
<td>Munich</td>
<td>✓</td>
<td>✓</td>
<td>Calibration to specific MR sequence</td>
<td>Required</td>
<td>1 m</td>
</tr>
<tr>
<td>CAR-RiDR</td>
<td>✓</td>
<td>✓</td>
<td>Calibration to specific MR sequence</td>
<td>Required</td>
<td>1 m</td>
</tr>
<tr>
<td>RESOLUTE</td>
<td>✓</td>
<td>✓</td>
<td>Calibration to specific MR sequence</td>
<td>Required</td>
<td>15 m</td>
</tr>
</tbody>
</table>
Engineering Sciences. Funding was also received from the National Institute for Health Research University College London Hospitals Biomedical Research Centre (NIHR BRC UCLH/UCL High Impact Initiative BW.mrn.BRC10269) and the EPSRC (EP/K005278/1). In Lyon, this work was supported by a CIFRE studentship jointly funded by Siemens and the French national agency for research and Technology (ANRT). Funding was also received from the French national ‘invest for the future’ programmes (LILLI – Lyon Integrated Life Imaging: hybrid MR-PET-ANR-11-EQPX-0026) and the hospital University Institute CESAME (Brain and Mental Health ANR-10-IBHU-0003). In Munich, this work was supported by the European Union Seventh Framework Program (FP7) under Grant agreement n° 602621 Trimage and n° 294582- MUMI. In Boston, this work was supported by the NIH National Institute of Biomedical Imaging and Bioengineering (NIBIB) grant number 1RO1EB014894.

Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.neuroimage.2016.12.010.

References