Low-dose CT for the spatial normalization of PET images: A validation procedure for amyloid-PET semi-quantification

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ABSTRACT

The reference standard for spatial normalization of brain positron emission tomography (PET) images involves structural Magnetic Resonance Imaging (MRI) data. However, the lack of such structural information is fairly common in clinical settings. This might lead to lack of proper image quantification and to evaluation based only on visual ratings, which does not allow research studies or clinical trials based on quantification.

PET/CT systems are widely available and CT normalization procedures need to be explored. Here we describe and validate a procedure for the spatial normalization of PET images based on the low-dose Computed Tomography (CT) images contextually acquired for attenuation correction in PET/CT systems. We included $N = 34$ subjects, spanning from cognitively normal to mild cognitive impairment and dementia, who underwent amyloid-PET/CT ($^{18}$F-Florbetaben) and structural MRI scans. The proposed pipeline is based on the SPM12 unified segmentation algorithm applied to low-dose CT images. The validation of the normalization pipeline focused on 1) statistical comparisons between regional and global $^{18}$F-Florbetaben-PET/CT standardized uptake value ratios (SUVrs) estimated from both CT-based and MRI-based normalized PET images (SUVr\textsubscript{CT}, SUVr\textsubscript{MRI}) and 2) estimation of the degrees of overlap between warped gray matter (GM) segmented maps derived from CT- and MRI-based spatial transformations.

We found negligible deviations between regional and global SUVrs in the two CT and MRI-based methods. SUVr\textsubscript{CT} and SUVr\textsubscript{MRI} global uptake scores showed negligible differences (mean ± sd 0.01 ± 0.03). Notably, the CT- and MRI-based warped GM maps showed excellent overlap (90% within 1 mm).

The proposed analysis pipeline, based on low-dose CT images, allows accurate spatial normalization and subsequent PET image quantification. A CT-based analytical pipeline could benefit both research and clinical practice, allowing the recruitment of larger samples and favoring clinical routine analysis.

1. Introduction

The evaluation of biomarkers for the early diagnosis of neurodegenerative conditions causing dementia has been increasingly recognized as of outmost importance in research and clinical practice (Ahmed et al., 2014; Albert et al., 2011; Armstrong et al., 2013; Dubois et al., 2014; Iaccarino et al., 2017; McKeith et al., 2017; McKhann et al., 2011a; Rascovsky et al., 2011; Sperling et al., 2011). As for Alzheimer's Disease (AD), the development of Positron Emission Tomography (PET) techniques to investigate brain amyloid accumulation (amyloid-PET) brought landmark changes in clinical neuroscience research (Villemagne, 2016). The reliability of PET tracers for in vivo amyloid assessment is supported by their correlation with post-mortem amyloid plaque measurement (Clark et al., 2012; Sabri et al., 2015; Wolk, 2011).

To date, their mandatory adoption in clinical trials and the great potential for diagnostic purposes is recognized, in particular to rule out AD pathology (Vandenberghe et al., 2013b; Vandenberghe et al., 2013a). Amyloid PET imaging plays a fundamental role for the inclusion and exclusion of subjects in clinical trials based on anti-amyloid treatments (Sperling et al., 2014a, 2014b), and it has been used as an
outcome measure as well (Salloway et al., 2014; Sevigny et al., 2016). The positivity of an amyloid-PET scan is commonly assessed qualitatively through a visual evaluation of the PET radiotracer distribution, in accordance to tracer-specific guidelines (Rowe and Vilmagne, 2013).

A correct and reliable quantification of regional amyloid burden with PET, however, is considered mandatory to avoid the limitations of the operator-dependent visual classification, especially in longitudinal studies (Perani et al., 2014a,b). The most commonly adopted semi-quantification techniques involve tracer-specific approaches to estimate regional amyloid burden based on Standardized Uptake Value Ratio (SUVR) measurements. SUVRs are obtained by comparing tracer uptake in target regions to a reference area devoid of specific uptake. By comparing SUVRs in AD patients with SUVRs obtained in healthy volunteers, previous studies derived cut-off thresholds that could discriminate between amyloid positive and amyloid negative individuals (Barthel et al., 2011; Chiotis et al., 2015; Fleisher, 2011; Nordberg et al., 2013; Oh et al., 2015; Ong et al., 2015; Vandenbergh et al., 2010). Semi-quantitative amyloid burden is generally estimated on average (composite) SUVR based on neocortical regions, usually including frontal, parietal, temporal and cingulate regions (Clark et al., 2012; Fleisher, 2011). The adopted reference regions for amyloid PET SUVR can vary and may include the whole cerebellum, the cerebellar gray matter (GM) and/or specific portions of the white matter (WM) (Brendel et al., 2015; Schmidt et al., 2015). Of note, the implementation of semi-quantification techniques can also introduce variability, especially with respect to differences in analysis procedures and scan protocols. All these factors can heavily impact the classification of amyloid burden, with considerable effects in research studies and consequences in clinical trials (e.g. inclusion/exclusion of subjects). Among the most important factors there are: i) the selection of regions of interest (ROIs); ii) the selection of reference regions and iii) the choice of running quantifications in either native or standard space, with the latter being strongly influenced by the spatial normalization algorithms.

In an ideal setting, structural Magnetic Resonance Imaging (MRI) scans are available for each subject, allowing high precision spatial normalization and ROIs definition. Conveying the PET images to standard space can offer the use of standardized, published atlases with regions of interest, such as Automatic Anatomical Labeling: AAL (Tzourio-Mazoyer et al., 2002), Talairach Daemon: TD (Lancaster et al., 2000; Lancaster et al., 1997), Individual Brain Atlas: IBA (Aleman-Gomez et al., 2006), allowing for the definition of ROIs and to estimate regional SUVRs. In a routine diagnostic setting, however, MRI data are not always available, thus preventing an MRI-based spatial normalization of the amyloid-PET images to a standard space. For many centers, the lack of an MRI-based normalization pipeline prevents any further quantification.

To overcome this limitation, several PET-only pipelines for spatial normalization have been developed, based on custom or simulated PET templates (Hutton et al., 2015; Lundqvist et al., 2013; Saint-Aubert et al., 2014). These templates enable an accurate PET image warping and semi-quantification, but they are tracer-specific, limiting their utilization to radioligands with similar radioactivity distributions. Furthermore, to perform an appropriate PET-based normalization, the tracer uptake should define brain anatomy in sufficient details, which is not always the case for PET molecular imaging radiotracers. Finally, the spatial distribution of the tracer should be reasonably similar across subjects, to prevent bias in registration. This is not the case for amyloid tracers, where tracer distribution varies markedly across individuals, depending on the degree of amyloid burden: while in positive cases GM uptake is on par with WM uptake, amyloid-negative subjects display high contrast between the two.

Building on these premises, there is a need for validated methods to perform reliable spatial normalization of PET amyloid images. In this view, and considering that most PET clinical studies are nowadays performed using PET/Computed Tomography (CT) systems, we tested and validated a method for a high precision spatial normalization and SUVr computation using the low-dose CT image acquired for attenuation correction (AC). The inclusion of a CT-based analytical pipeline for PET quantification would allow a net benefit in terms of both research and clinical practice, allowing the recruitment of larger samples and favoring clinical routine analysis.

2. Materials and methods

2.1. Participants

Subjects were retrieved from the Ricerca Finalizzata Progetto di Rete Nazionale AD (AD-NETWORK/RETEAD) database. RETEAD is a large Italian multicenter study that aims at developing and validating operational research criteria for diagnosis of AD in the preclinical/predementia phase and early recognition of atypical forms, based on a multi-factorial protocol that integrates molecular, imaging, neuropsychological and clinical profiles. The study conformed to the ethical standards of the Declaration of Helsinki for protection of human subjects. Each subject provided written informed consent as approved by the Local Ethical Committees.

Thirty four subjects (age = 69.58 ± 6.63 (range:50–80) years; M/F = 16/18) were recruited at Fondazione IRCCS Istituto Neurologico Besta, Milan. The sample consisted of subjects in preclinical and prodromal dementia phases and patients with overt dementia, thus covering a wide spectrum of cases, from normal cognition to dementia. In detail, the sample included 4 subjects with subjective cognitive complaints (Jessen et al., 2014), 12 subjects with pre-mild cognitive impairment (pre-MCI) (Storandt et al., 2006), 14 subjects with MCI (8 single-domain MCI and 6 multi-domain MCI) (McKhann et al., 2011b) and 4 patients with a diagnosis of probable AD dementia (McKhann et al., 2011a).

Each subject underwent brain structural imaging, including an MRI scan at Fondazione IRCCS Istituto Neurologico Besta, Milan and an amyloid PET/CT scan at the Nuclear Medicine Unit of San Raffaele Hospital, Milan. Inter-scan interval was no longer than six months for MRI and amyloid PET scans.

2.2. Image acquisition

2.2.1. 18F-Florbetaben PET/CT

Each subject received an intravenous injection of 300 ± 37 MBq of 18F-Florbetaben (Neuracq, Piramal). The dose was administered as a single bolus injection followed by 20 cc of saline flush. All PET acquisition were performed using a hybrid PET/CT Discovery-690 system (General Electric Medical Systems Milwaukee, WI, USA) (Bettinardi et al., 2011). After positioning, a low dose CT scan (kVP: 140 kV, current: 40 mA, rotation time: 0.8 s, slice thickness: 3.75 mm, pitch: 1.375:1) was acquired to be used for attenuation correction of PET data. Images were reconstructed using the “standard” kernel, a 30 cm reconstruction field of view, and 3.27 mm slice interval, for a resulting voxel size of 0.59 × 0.59 × 3.27 mm³. A 3D-PET acquisition (list mode) was started about 90 min after the injection of the tracer and lasted for 20 min. Image reconstruction was performed by using a 3D Ordered Subsets Expectation Maximization (OSEM) algorithm with the following parameters: Image matrix = 128, Field Of View = 250 mm, Subsets = 24, Iterations = 3, Post Filter (Gaussian) = 3 mm FWHM, Attenuation Correction = CT-based. The resulting voxel size was 1.95 × 1.95 × 3.27 mm³.

2.2.2. MRI

The MRI imaging data were acquired in Neurological Institute “C. Besta”, using an Achieva 3 T MR scanner (Philips Healthcare BV, Best, NL) equipped with a 32-channel head coil. A volumetric turbo field echo (TFE) T1-weighted structural sequence (180 sagittal slices, TR = 8.3 ms, TE = 3.9 ms, FOV = 240 × 240 mm, voxel size = 1x1x1 mm³, flip angle = 8°) was acquired for each subject. Other structural, diffusion and functional magnetic imaging data were also
collected in the same session, but not reported in this study. The total duration of scanning session was around 55 min.

2.2.3. Image processing

All CT and MRI images were converted from the original DICOM to a NIFTI format using SPM12. The origin coordinates of the MRI scan were manually set to the anterior commissure. CT and PET images were rigidly co-registered to the respective MRI scan. A visual inspection was always performed as a quality control to detect possible errors in the co-registration step.

2.3. Spatial normalization

2.3.1. CT-based normalization algorithm

The hypothesis of the proposed procedure is that a low dose CT scan contains enough information in terms of contrast between GM, WM and cerebro-spinal fluid (CSF) to estimate the spatial transformation accurately. To spatially normalize the CT images into the stereotactic standard space, we optimized the unified segmentation-normalization algorithm as implemented by Ashburner and Friston in SPM12 (Ashburner and Friston, 2005). This algorithm iteratively finds the spatial transformation that best matches the aligned images to a set of tissue probability maps (TPM), that are used in a subsequent tissue classification procedure.

The algorithm works with a parametrization of the image intensities for each tissue type using a Gaussian Mixture Model (GMM). In the latest version, known as “new segment”, included in SPM12 (Malone et al., 2015; Weiskopf et al., 2011), 6 tissue classes are considered: GM, WM, CSF, bone, outside tissues and air. A priori TPM are used to improve the segmentation, to fix initial intensities for each tissue class and to distinguish tissues with identical average intensities. This iterative algorithm first estimates the spatial deformation from the Montreal Neurological Institute (MNI) space to the subject native space. The TPM are then transformed to the subject native space using this deformation and they are used to update the GMM describing the intensities of all tissues. Finally, the spatial transformation from the MNI space to the native space is updated, looking for the one that best matches the current GMM. The procedure is iterated until the algorithm converges. Forward and backward transformations are both estimated.

In our proposed optimization, the CT images are pre-processed by a “clean-up” procedure. Every value lower than −300 HU is set to −1024 HU, to avoid low-density structures outside the head (e.g.: head-holder, cushion etc.) confounding the algorithm. Without this procedure, the coarse affine registration performed before the actual segmentation often fails. The resulting images are then loaded in the SPM12 unified segmentation algorithm, with optimized parameters. Compared to the default SPM12 settings for MRI, we disabled bias-field correction, as CT does not suffer from this artifact. Also, as Hounsfield values are generally comprised in a broad range in CT, we used only 1 Gaussian for the GMM of GM, WM, and only 2 Gaussians for the other 4 classes (CSF, bone, outside tissue and air). Identical regularization strengths to the default were used. All the parameters used are summarized in Table 1. The number of Gaussians was chosen a priori, knowing that in CT GM and WM have only one average intensity. For CSF, 2 Gaussians were used to take into account the possibility that CSF close to the bone has higher intensity than that in the ventricles due to spillover effect. As bone has values in a very broad range in CT, 2 gaussians were used. Tissue can be thought as composed of low-intensity fat and higher-intensity areas like muscles, so 2 gaussians also were used. In the “air” class we expect all low intensity structures that have the same value of −1024, after our thresholding, modeled with 1 gaussian, and another Gaussian to model all the other low intensity pixels between about −50 to −300 HU.

The robustness of our algorithm to different settings was tested in supplementary material.

2.3.2. MRI-based spatial normalization

MRI images were spatially normalized to the MNI space using the SPM12 unified segmentation and default settings. The parameters are reported in Table 1, to be compared to the optimization introduced for the CT version. The segmented tissues and the forward transformation to the MNI space are then saved. The spatial transformation was used as a comparison with the CT one. The GM map was also used in a later stage to compare and validate the proposed method (see below).

2.4. Comparison of the CT and MRI normalizations

Two strategies were used to compare the output of the normalization procedures, with the MRI-based normalization considered as the gold standard for the pre-processing step. First, we measured, from the two normalized CT (nPET CT) and MRI-based (nPETMRI) 18F-Florbetaben PET images, the SUVr values in a set of ROIs commonly considered to estimate amyloid burden in AD. The second strategy consisted in measuring, for each subject, the degree of overlap between the two estimated GM maps (GMCT and GMMRI) obtained with the CT- and the MRI-derived transformations. Normalized 18F-Florbetaben PET images were resampled to a bounding box of [−90–126 -72; 90 90,108] mm with an isotropic voxel size of 2 mm, using the two previously estimated deformations.

2.4.1. 18F-Florbetaben SUVr estimation

Multiple ROIs for regional and global amyloid burden assessment were selected according to previous literature (Barthel et al., 2011; Ong et al., 2013; Sabri et al., 2015; Tiepolt et al., 2016) To summarize, we extracted six ROIs representing wide, bilateral cortical macroareas (Fig. 1), i.e. Dorsolateral and Medial Frontal Cortex, Cingulum, Precuneus, Inferior and Superior Parietal Lobules, Lateral Occipital Cortex, and Lateral Temporal Cortex, from the Automated Anatomical Labeling (AAL) atlas (Tzourio-Mazoyer et al., 2002), through the Wake Forest University PickAtlas toolbox for SPM (Maldjian et al., 2003). All the images were scaled to the activity of the cerebellar GM, used as the reference region (Catafau et al., 2016; Villemagne et al., 2015). The average computed from the six ROIs was considered as an index of global cortical amyloid burden.

2.4.2. Statistical analysis

Concordance between the SUVr measures was assessed using Bland-Altman plots and estimation of region-wise Pearson correlation analysis. Regional and global amyloid burdens and standard deviations were computed with mean absolute differences and limits of agreement (see Table 2).
voxels in the GMCT that overlap with those in the GMMRI. It is reasonable to expect a complete overlap of the two maps in the case of SUVrMRI (mean ± sd 0.012 ± 0.032), as further con-

There was a trend with −

Regional SUVr computed on the whole brain considering the two different spatial normalization pipelines (see text). Values are shown as means, while limits of agreement are computed as: mean(d)-1.96’sd(d) | mean(d) + 1.96’sd(d).

2.4.3. Comparison of gray matter maps
The GM maps obtained from the MRI segmentation of each patient were resampled to the MNI space within a bounding box of [−90−126−72; 90, 90, 108] with an isotropic voxel size of 1 mm, to better capture the limited thickness of GM. The MRI and the CT derived deformations were used to obtain a GMMRI and a GMCT map for each subject. It is reasonable to expect a complete overlap of the two maps in the case of identical deformations. Therefore, we measured the percentage of voxels in the GMCT that overlap with those in the GMMRI. The GMMRI was then dilated to the 6 nearest-neighbor voxels in three dimensions, and this dilated map was used again as a reference to perform the same computation. This allowed the assessment of the fraction of GM that was transformed either to the correct location or to a 1 mm wide neighborhood. This dilation was performed two more times to measure concordance within 2 mm and 3 mm wide from the reference. These overlap measures were computed over the whole image and also locally, inside the same ROIs defined for the SUVr analysis.

3. Results
3.1. 18F-Florbetaben SUVr
The main steps of the proposed procedure are shown in Fig. 2. The mean regional SUVrCT and SUVrMRI as well as the composite cortical values are reported in Table 2. Negligible differences were found at regional level. Occipital SUVr scores showed some deviation, but were still negligible (mean absolute difference 0.048, limits of agreement −0.067 | 0.160), whereas the temporal SUVr scores showed the best concordance (mean absolute difference 0.010, limits of agreement −0.079 | 0.056). When compared to the regional SUVrMRI, the SUVrCT was slightly underestimated for all regions, except for the occipital region which showed the opposite trend. Global amyloid burden showed very narrow differences using average regional SUVrCT or SUVrMRI (mean ± sd 0.012 ± 0.032), as further confirmed by the Bland-Altman plot (Fig. 3) and correlation analysis (Pearson r = 0.994, p < 2.2e-16) (Fig. 3). In the Bland-Altman plot of the global SUVr, we performed a Spearman correlation to test whether there was a trend between the SUVrCT vs SUVrMRI difference and the global SUVr load. There was a trend with $r^2 = 0.16$, which is significant with $p = .02$.

In supplementary materials we show that these results are affected by the exact algorithms settings, at least when using reasonable parameters.

3.2. Comparison of gray matter maps
The overlap between GM maps obtained using the two different transformations is shown in Fig. 4. Transforming the GM maps using the CT-based normalization algorithm provided a 70% overlap with those obtained with MR-based normalization. Notably, > 90% of the GM map voxels from the CT-based normalization were within 1 mm from the MR-transformed ones. Full results including 2 and 3 mm dilation are reported in Table 3.

4. Discussion
A correct AD diagnosis in early prodromal, and even preclinical, disease stages is important, especially for the appropriate inclusion of patients in clinical trials (Sperling et al., 2014a, 2014b). In this framework, amyloid-PET is unique for the detection of pathological Aβ accumulation in vivo (Vandenberghe et al., 2013a,b). Amyloid positivity, as shown by PET studies, is currently considered as a supportive biomarker for AD diagnosis according to the diagnostic criteria (Albert et al., 2011; Dubois et al., 2014; McKhann et al., 2011a; Sperling et al., 2011). Validated and standardized amyloid (semi)quantification procedures are therefore needed to overcome the limitations of visual ratings and/or binary classifications, for both diagnostic and research purposes (Perani et al., 2014), and also for a better evaluation of cases for clinical trials. To this end, there is a need to develop highly reliable (semi)quantification approaches to accurately measure amyloid burden in different brain regions at the single-subject level. Previous studies have focused on how to optimize Amyloid-PET analysis to improve both quantification (Bullich et al., 2017a; Saint-Aubert et al., 2014) and reliability of longitudinal PET assessments (Brendel et al., 2015; Bullich et al., 2017b). Notably, semi-quantification with SUVr increased the classification accuracy in comparison to visual ratings (Bullich et al., 2017a; Camus et al., 2012; Perani et al., 2014b).

The majority of previous studies has focused on the development of highly accurate processing steps for the quantification, especially considering reference and target region selection and definition of optimal scanning protocols. The present study, however, focuses on the pre-processing phase, with the aim of validating a feasible CT-based pipeline which could greatly increase the implementation of semi-quantification protocols in research and clinical settings.

When performing PET data analysis in standard space, spatial normalization is the first and a crucial step for the accurate estimation of radioligand specific uptake. High-precision spatial alignment of brain structures is indeed fundamental to achieve the highest statistical power. High resolution MRI images are currently considered the gold standard for spatial warping and many algorithms are available to compute the individual spatial transformations. MRI scans require extra costs and time, on top of being a procedure that cannot be performed on all patients (e.g. in presence of metallic inserts, or due to claustrophobia). While these limitations might not be relevant in research settings, MRI scans are not always acquired in routine studies in clinical settings.

![Fig. 1. Visualization of the regions of interest used for the analysis, overlaid on a standardized template.](image-url)
The CT acquisition protocol resulted in a volumetric CT Dose Index (CTDIvol) of about 1.5 mGy, which translated to a Dose Length Product (DLP) of about 25 mGy*cm, meaning, in an average patient, a dose of < 70 μSv. This dose provides sufficient anatomical details while being extremely low. In this work CT were acquired using 40 mA current and 0.8 s rotation time. The most recent scanners, like ours, can be set up to acquire images with currents as low as 10 mA and lower rotation times, for even lower doses. However, not only the image quality significantly decreases, but also there is the risk of "photon starvation" artefacts, which lead to incorrect attenuation correction (Xia et al., 2012). To overcome this limit, new methods are being used where the current is not lowered but the number of projections acquired is reduced, using then sparse view reconstruction techniques (Rui et al., 2015). However, these techniques have only been recently introduced clinically and they have not been optimized to provide anatomical detail.

A limit of this study is that the influence of CT image quality on the algorithm could not be studied. Future studies will need to assess both the lower dose limit at which the algorithm still performs correctly, and also whether the use of diagnostic quality CT provides improvements.

Here we found high concordance with MRI-based normalization using the CT-based normalization procedure. Notably, considering the SUVr values, the standard deviation of the differences between the reference MRI-based normalization and the proposed algorithm was < 0.05 in all the considered ROIs. Therefore, this semi-quantitative approach allowed for an accurate measurement of the amyloid burden in each ROI, which is of importance since regional variations of amyloid burden are well-documented in literature (Jansen et al., 2015; Ossenkoppele et al., 2015). Additionally, in the second validation test, comparing the CT- and MRI-based GM maps registration, we found that the maps overlapped within 1 mm in 90% of the voxels. This validation was run on a patient population that spanned over a wide range of pathological stages and of amyloid burden, where brain anatomy might differ due to increased atrophy. Accordingly, global amyloid SUVr for our population ranged from 0.9 to 1.8 SUVr. A very small trend, $\rho^2 = 0.16$, was noted in the Bland-Altman plot. Considering that the $p$-value was at the threshold for statistical significance ($p = .02$), this result might be due to the presence of few outliers. Studying this algorithm on larger number of subjects, especially AD with high amyloid load and high levels of atrophy, could further certify the good
The performance of this algorithm in all the possible conditions. In Supplementary Materials, we have also shown that the algorithm is robust to changes in the settings, concerning to how the thresholding operation is performed and to the number of gaussians used for outside tissues. The proposed CT normalization procedure was developed for hybrid PET/CT scanners, since this is the most widely diffuse system for both clinical and research applications. This approach does not require any tracer-specific template to determine the spatial normalization. Notably, while the present procedure was validated using the \(^{18}\)F-Florbetaben tracer, it can be adopted as well with other radioligands. This CT-based approach could also be useful for analyzing large retrospective databases where CT, but not MRI, scans are only available. Another important application could be the development of an automated processing pipeline that can aid clinicians in the evaluation of amyloid PET scans. Finally, it should be noted that this procedure was built exclusively using widely available and validated free tools: SPM and the AAL atlas.

5. Conclusions

In this work, we validated an automated spatial normalization method for amyloid PET data, based on the low-dose CT acquired contextually with the PET scan for attenuation correction. This procedure, when compared to the gold-standard MRI-based spatial transformation, showed extremely high levels of concordance in the results of semi-quantification. The procedure is well-suited for both clinical and research applications as well as for radioligands other than amyloid-tracers since it has a simple implementation and is based on validated and widely available tools.

Conflict of interest

The authors declare that they have no conflicts of interest regarding the publication of this article.

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Declarations of interest

None.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.nicl.2018.07.013.

References

Ahmed, R.M., Paterson, R.W., Warren, J.D., Zetterberg, H., O’Brien, J.T., Fox, N.C., Halliday, G.M., Schott, J.M., 2014. Biomarkers in dementia: clinical utility and new directions. J. Neurol. Neurosurg. Psychiatry 85, 1426-1434. https://doi.org/10.1136/jnnp-2014-307662.
Catafau, A.M., Bullich, S., Seibyl, J.P., Barthel, H., Ghetti, B., Leverenz, J., Ironside, J.W., Bullich, S., Seibyl, J., Catafau, A.M., Jovalekic, A., Koglin, N., Barthel, H., Sabri, O., De Iaccarino, L., Sala, A., Caminiti, S.P., Perani, D., 2017. The emerging role of PET imaging in dementia. Alzheimers & Dementia 7, 270–279. https://doi.org/10.1016/j.jalz.2016.03.008.

Aleman-Gomez, Y., Melie-Garcia, L., Valdés-Hernandez, P., 2006. IBASPM: toolbox for expert-aided detection of cortical atrophy. NeuroImage 32, 165–171. https://doi.org/10.1016/j.neuroimage.2006.03.044.

Guilloteau, D., 2012. Using PET with 18F-AV-45 (florbetaben Aβ) in patients with Alzheimer’s disease and healthy controls: a multicentre 2 diagnosis study. Lancet Neurol. 11, 424–435. https://doi.org/10.1016/S1474-4422(11)70077-1.

Bettinardi, V., Presotto, L., Rapisarda, E., Picchio, M., Giondolo, L., Gilardi, M.C., 2011. Physical performance of the new hybrid PET/CT Discovery-690. Med. Phys. 38, 5394–5411. https://doi.org/10.1118/1.3635220.

Brendel, M., Högenauer, M., Deller, A., Sauerbeck, J., Barthel, P., Seibyl, J., Rüther, M., 2014. Improved longitudinal [18F]AV-45 amyloid PET by white matter reference and VOI-based partial volume effect correction. NeuroImage 108, 450–459. https://doi.org/10.1016/j.neuroimage.2014.11.055.

Bullich, S., Seibyl, J., Catafau, A.M., Jovalekic, A., Koglin, N., Barthel, H., Sabri, O., De Santi, S., 2017a. Optimized classification of 18F-FITBETAPET scans as positive and negative using an SQRV quantitative approach and comparison to visual assessment. NeuroImage 151, 325–332. https://doi.org/10.1016/j.neuroimage.2017.04.025.

Bullich, S., Villemagne, V.L., Catafau, A.M., Jovalekic, A., Koglin, N., Rowe, C.C., De Santi, S., 2017b. Optimal reference region to measure longitudinal amyloid-β change with 18F-FITBETAPET. J. Nucl. Med. 58, 1300–1306. https://doi.org/10.2967/jnumed.116.178351.

Camus, V., Payoux, P., Barré, L., Desgranges, B., Voisin, T., Tauber, C., La Joie, R., Tafani, M., Hommet, C., Chételat, G., Mondon, K., de La Sayette, V., Cottier, J.P., Beauclerc, J., Salmon, D., Thirion, B., 2015. Cortical atrophy in healthy aging. NeuroImage 108, 450–459. https://doi.org/10.1016/j.neuroimage.2014.11.055.

Catafau, A.M., Bullich, S., Seibyl, J., Barthel, H., Ghetti, B., Leverenz, J., Iwamoto, J., Schulz-Schaeffer, W.J., Hoffmann, A., Sabri, O., 2016. Cerebellar amyloid-β plaques: how frequent are they, and do they influence 18F-FITBETAPET SUV ratios? J. Nucl. Med. 57, 1740–1745. https://doi.org/10.2967/jnumed.116.171562.

Choiti, K., Carter, S.P., Farid, K., Savitcheva, I., Nordberg, A., 2015. Amyloid PET in European and North American cohorts; and exploring age as a limit to clinical use of florbetaben Aβ imaging. J. Nucl. Med. 56, 1300–1306. https://doi.org/10.2967/jnumed.116.178351.

Clark, M.G., Pontecorvo, M.J., Beach, T.G., Bedell, B.J., Coleman, R.E., Douraswavith, P.M., Fleisher, A.S., Reiman, E.M., Sabbagh, M.N., Sadowsky, C.H., Schneider, J.A., Arora, A., Carpenter, A.P., Flitter, M.L., Joshi, A.D., Krautkramer, M.J., Lu, M., Minton, M., Peskind, E.R., Pontecorvo, M.J., 2012. Cerebral PET imaging in atypical parkinsonism at autopsy for tauopathology at autopsy for detection of neuritic amyloid-β plaques: a prospective cohort study. Lancet Neurol. 11, 669–678. https://doi.org/10.1016/S1474-4422(12)70142-4.

Dubois, B., Feldman, H.H., Jacova, C., Hampel, H., Molinuevo, J.L., Scheltens, P., Sluimer, J., Zetterberg, H., 2014. Criteria for the diagnosis of corticobasal degeneration. Neurology 80, 496–503. https://doi.org/10.1212/WNL.0000000000000631.

Dickson, D.W., Grossman, M., Hallett, M., Josephs, K.A., Kertesz, A., Lee, S.E., Miller, D.J., van Buchem, M.A., Camus, V., Cavedo, E., Chen, K., Chetelat, G., Cohen, A.D., Dubois, B., Feldman, H.H., Jacova, C., Hampel, H., Molinuevo, J.L., Scheltens, P., Sluimer, J., Zetterberg, H., 2014. Criteria for the diagnosis of corticobasal degeneration. Neurology 80, 496–503. https://doi.org/10.1212/WNL.0000000000000631.

Dubois, B., Feldman, H.H., Jacova, C., Hampel, H., Molinuevo, J.L., Blennow, K., Dekosky, S.T., Gauthier, S., Selkoe, D.J., van Buchem, M.A., Skovronsky, D.M., 2012. Cerebral PET with 18F-Florbetaben for the detection of cerebral Aβ plaques: a prospective multicentre PET study of fibrillar amyloid in Alzheimer’s disease. Eur. J. Nucl. Med. Mol. Imaging 40, 104–112. https://doi.org/10.1007/s00259-012-2118-9.

Dubois, B., Feldman, H.H., Jacova, C., Hampel, H., Molinuevo, J.L., Blennow, K., Dekosky, S.T., Gauthier, S., Selkoe, D.J., van Buchem, M.A., Skovronsky, D.M., 2012. Cerebral PET with 18F-Florbetaben for the detection of cerebral Aβ plaques: a prospective multicentre PET study of fibrillar amyloid in Alzheimer’s disease. Eur. J. Nucl. Med. Mol. Imaging 40, 104–112. https://doi.org/10.1007/s00259-012-2118-9.

Hutton, C., Declerck, J., Mintun, M.A., Nutt, J.G., Sutra, T., Phelps, C.H., 2011b. The diagnosis of dementia due to Alzheimer’s disease: recommendations from the National Institute on Aging-Alzheimer’s Association workgroups on diagnostic guidelines for Alzheimer’s disease. Alzheimers & Dementia 7, 263–269. https://doi.org/10.1016/j.jalz.2011.03.005.

McKhan, G.M., Knopman, D.S., Cherokoff, H., Hyman, B.T., Jack, C.R., Kawas, C.H., Klunk, W.E., Koroszetz, W.J., Manly, J.J., Mayeux, R., Mohs, R.C., Morris, J.C., Rossor, M.N., Scheltens, P., Carrillo, M.C., Thies, B., Weintraub, S., Phelps, C.H., 2011a. The diagnosis of mild cognitive impairment due to Alzheimer’s disease: recommendations from the National Institute on Aging-Alzheimer’s Association workgroups on diagnostic guidelines for Alzheimer’s disease. Alzheimers & Dementia 7, 263–269. https://doi.org/10.1016/j.jalz.2011.03.005.

Jessen, F., Amarniello, R.E., van Boxstel, M., Bretert, M., Ceccaldi, M., Chételat, G., Dubois, B., De Deyn, P.P., Frisoni, G.B., Forno, D., Freytag, S., Garibotto, V., Almkvist, O., Kalbe, E., Hinz, R., Herholz, K., 2013. A European multicentre PET study of fibrillar amyloid in Alzheimer’s disease. Eur. J. Nucl. Med. Mol. Imaging 40, 104–114. https://doi.org/10.1007/s00259-012-2237-2.

Oh, H., Steffener, J., Radligh, Q.R., Habek, C., Liu, D., Gazes, Y., Janicki, S., Sten, Y., 2015. MRI-based hyperperfusion to improve the identification of cognitively normal elderly. Neurobiol. Aging 36, 3247–3254. https://doi.org/10.1016/j.neurobiolaging.2015.08.016.

Koh, K.Y., Villemagne, V.L., Bahar-Fuchs, A., Lamb, P., Chételat, G., Raniga, P., Pallu, M., Salvado, O., Putz, B., Roth, K., Masters, C.L., Reinchinger, B., Rowe, C.C., 2013. 18F-florbetaben Aβ imaging in mild cognitive impairment. Alzheimers Res. Ther. 5, 19. https://doi.org/10.1186/1719-5151-5-19.
