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Original research

Added value of FDG-PET for detection of progressive supranuclear palsy

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ABSTRACT

Background Diagnostic criteria for progressive supranuclear palsy (PSP) include midbrain atrophy in MRI and hypometabolism in [¹⁸F]fluorodeoxyglucose (FDG)-positron emission tomography (PET) as supportive features. Due to limited data regarding their relative and sequential value, there is no recommendation for an algorithm to combine both modalities to increase diagnostic accuracy. This study evaluated the added value of sequential imaging using state-of-the-art methods to analyse the images regarding PSP features.

Methods The retrospective study included 41 PSP patients, 21 with Richardson's syndrome (PSP-RS), 20 with variant PSP phenotypes (vPSP) and 46 sex- and age-matched healthy controls. A pretrained support vector machine (SVM) for the classification of atrophy profiles from automatic MRI volumetry was used to analyse T1w-MRI (output: MRI-SVM-PSP score). Covariance pattern analysis was applied to compute the expression of a predefined PSP-related pattern in FDG-PET (output: PET-PSRP expression score).

Results The area under the receiver operating characteristic curve for the detection of PSP did not differ between MRI-SVM-PSP and PET-PSRP expression score ($p \geq 0.63$): about 0.90, 0.95 and 0.85 for detection of all PSP, PSP-RS and vPSP. The MRI-SVM-PSP score achieved about 13% higher specificity and about 15% lower sensitivity than the PET-PSRP expression score. Decision tree models selected the MRI-SVM-PSP score for the first branching and the PET-PSRP expression score for a second split of the subgroup with normal MRI-SVM-PSP score, both in the whole sample and when restricted to PSP-RS or vPSP.

Conclusions FDG-PET provides added value for PSP-suspected patients with normal/inconclusive T1w-MRI, regardless of PSP phenotype and the methods to analyse the images for PSP-typical features.

INTRODUCTION

Progressive supranuclear palsy (PSP) is a primary 4-repeat tauopathy^{1,2} presenting with a broad

WHAT IS ALREADY KNOWN ON THIS TOPIC

⇒ Current diagnostic criteria list midbrain atrophy on structural MRI and hypometabolism on brain [¹⁸F]fluorodeoxyglucose (FDG)-positron emission tomography (PET) as supportive progressive supranuclear palsy (PSP) features, but due to the lack of data, they do not make any recommendations on the sequential use of both imaging modalities in patients where the clinical status and one imaging modality remain inconclusive. This contributes to the lack of consensus on how neuroimaging should be used in the diagnosis of PSP.

WHAT THIS STUDY ADDS

⇒ This study indicates that brain FDG-PET is about 15% more sensitive for the detection of PSP than structural MRI and, therefore, might be clinically useful for the detection of PSP in patients with normal or inconclusive structural MRI, irrespective of the PSP predominance type.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

⇒ These findings can guide decision-making on sequential MRI and FDG-PET imaging in clinically unclear patients with suspected PSP.

phenotypic spectrum including the 'classical' Richardson syndrome (PSP-RS) and various other variants (vPSP).^{3,4} Despite the fact that accounting for the large phenotypic variability in the revised criteria of the PSP study group of the International Parkinson and Movement Disorder Society (MDS) improved the PSP diagnosis, particularly at early disease stages and for vPSP,^{3,5} there is a need for supportive features to assist the clinical diagnosis in uncertain cases.^{6,7} The MDS criteria list midbrain atrophy on structural MRI and hypometabolism on [¹⁸F]

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fluorodeoxyglucose (FDG) positron emission tomography (PET) of the brain as supportive PSP features.^{3,8}

There are numerous studies on either structural MRI or FDG-PET alone for the (differential) diagnosis of PSP.^{8–19} However, there is a striking lack of head-to-head comparisons of both modalities in the same patients. In particular, there are no studies on sequential administration of both modalities in patients where the clinical status and one imaging modality remain inconclusive. This contributes to the lack of consensus on how neuroimaging should be used in the diagnosis of parkinsonian syndromes.¹⁶

Structural MRI and FDG-PET can be assessed visually for descriptive PSP features/signs. This can be supported by quantitative and/or statistical analyses in predefined regions-of-interest or voxel-by-voxel. However, recent studies demonstrated that fully automatic multivariable analyses using a support vector machine (SVM) to classify volumetric MRI profiles and covariance analysis of FDG uptake patterns clearly outperform visual analyses (with and without support by univariate quantitative/statistical analyses) in the detection of PSP, most pronounced in vPSP.^{10,18}

The current multicentre study performed a head-to-head comparison between SVM classification of MRI atrophy profiles and covariance pattern analysis of FDG-PET in the same clinical sample including about 50% vPSP. The a priori hypothesis was that covariance pattern analysis of FDG-PET is more sensitive for the detection of PSP than SVM classification of MRI atrophy profiles so that additional FDG-PET is most useful in case of normal or inconclusive MRI.

METHODS

Subjects

This retrospective multicentre study was designed as phase 2 study ('ability to discriminate patients from controls') according to the five-phase framework for biomarker validation²⁰ and, therefore, included well-characterised patients with established PSP diagnosis and healthy controls (HCs).

The archives of the university hospitals of Augsburg, Hamburg, Hannover, Leipzig and Munich were searched for consecutive patients using the following inclusion criteria: (1) clinical PSP diagnosis according to the MDS criteria by a movement disorder specialist and (2) structural MRI and FDG-PET performed within 12 months. 63 patients fulfilling these criteria were identified. From these, 18 patients (28.6%) were excluded because the T1w sequence was not adequate for morphometric/volumetric analyses, 4 patients (6.3%) because of severe ischaemic small vessel disease (Fazekas grade 3).²¹ The remaining 41 patients were included in the analyses (age at PET 69.6±7.8 years, range 53–81 years, 49% females). Mean disease duration at PET was 3.1±2.6 years (range 0.4–12.2 years). Median and IQR of the delay between the date of the clinical reference diagnosis and FDG-PET were 0 day and –6 days to 6 days, respectively (range –210 to 441 days).

Diagnostic certainty was probable/possible/suggestive of PSP in 35 (85.4%)/1 (2.4%)/5 (12.2%) patients. PSP-RS was the most frequent predominance type: n=21 (51.2%). Among the 20 patients with vPSP, 11 presented with a 'cortical' syndrome²²: 10 with predominant corticobasal syndrome (24.4% of all PSP patients) and 1 with predominant frontal presentation (2.4%). The remaining nine vPSP patients presented with a 'subcortical' syndrome: eight with predominant parkinsonism (19.5% of all PSP patients) and one with progressive gait freezing (2.4%). The patients scored 30±14 (range 9–86) on the PSP rating scale²³ and 21.4±4.4 (n=32, range 10–29) on the Montreal Cognitive

Assessment scale. Depressive symptoms scored 5.7±2.6 (n=28, range 1–12) on the 15-point Geriatric Depression Scale.

46 HC subjects from the Alzheimer's Disease Neuroimaging Initiative (ADNI-1) were included (age at PET 72.2±4.7 years, range 59–79 years, 39% females). The HC subjects did not differ significantly from the PSP patients with respect to age (p=0.07) or sex (p=0.37). The ADNI was launched in 2003 as a public-private partnership, led by principal investigator Michael W Weiner. For up-to-date information, see www.adni-info.org.

The same subjects had been included in previous studies on imaging-based detection of PSP with T1w-MRI¹⁸ or FDG-PET.¹⁰ These previous studies compared different approaches to identify PSP features separately in MRI and FDG-PET. In T1w-MRI, fully automatic classification of atrophy profiles by an SVM clearly outperformed descriptive signs (hummingbird, morning glory, Mickey-Mouse), visual reading supported by manual and automatic planimetric measures as well as visual reading supported by automatic volumetry.¹⁸ In FDG-PET, automatic covariance pattern analysis clearly outperformed visual interpretation of FDG-PET supported by voxel-based statistical testing.¹⁰ The current study first compared optimal (SVM) evaluation of T1w-MRI and optimal (covariance expression) evaluation of FDG-PET for the detection of PSP. Then, the putative added value of a sequential administration of these modalities was evaluated.

MRI

MRI was performed at 3/1.5 Tesla in 28/13 PSP patients (68.3/31.7%) with 15 different MR systems of 3 manufacturers (Siemens, Philips, Toshiba).¹⁸

For the ADNI HC subjects, the first of the two back-to-back 3D T1w-MRI from the baseline session was used.²⁴

SVM classification of T1w-MRI

For the automatic classification of T1w-MRI as PSP or HC, an SVM was used that had been trained previously using an independent dataset consisting of 106 PSP patients and 73 HCs.¹¹ The input to the SVM comprised the results of automated atlas-based MRI volumetry for 44 different compartments, brain structures, and planes, all corrected for intracranial volume and age and scaled to a range of 0–1 (online supplemental figure 1A). The output of the SVM consisted of an MRI-SVM-PSP score ranging between 0 (most likely HC) and 1 (most likely PSP). The pretrained SVM of 2016 was used without any changes.

FDG-PET imaging

In the PSP patients, brain FDG-PET was performed with six different PET systems (five PET/CT and one PET/MRI) from two manufacturers (Siemens, Philips) after intravenous injection of 202±37 MBq (range 128–317 MBq) according to the local standard operating procedure at each site. PET and MRI had been acquired simultaneously with a PET/MRI system in five patients. In the remaining 36 patients, median and IQR of the delay (absolute value) between PET and MRI were 7 days and 2–99 days, respectively. MRI had been performed before PET in 21 of the 36 patients (58.3%). The PET images were harmonised with respect to spatial resolution (8 mm full-width-at-half-maximum) as described previously.¹⁰

For the ADNI HC subjects, the baseline FDG-PET was used, fully preprocessed by the ADNI imaging core lab including filtering to 8 mm full width at half maximum.²⁵

PET images with harmonised resolution were spatially normalised to the Montreal Neurological Institute space using

the Statistical Parametric Mapping software package (SPM12) with default parameter settings.²⁶ The spatially normalised images were smoothed to spatial resolution of 12 mm full width at half maximum. No intensity scaling was performed (not required for covariance pattern analysis).

Covariance pattern analysis of FDG-PET

Spatial covariance analysis of FDG-PET was performed with scaled subprofile model principal component analysis^{27–29} implemented in the Scan Analysis and Visualization Processor software package (ScAnVP7.0w)^{28–30} freely available from the Feinstein Institute for Medical Research (<https://feinsteinneuropsychology.org/imaging-software/download-software>). This software was used for automatic computation of the expression of the predefined ‘North American’ PSP-related pattern (PSPRP)³¹ in each individual FDG-PET. The ‘North American’ PSPRP was derived in an independent sample of 10 patients with PSP diagnosis according to the NINDS criteria³² and 10 age- and sex-matched healthy volunteers (online supplemental figure 1B).³¹ No attempt was made to adapt the ‘North American’ PSPRP to the current dataset. The PET-PSPRP expression scores were corrected for age as described previously.¹⁰

Statistical analyses

MRI-SVM-PSP and PET-PSPRP expression score were transformed to z-scores relative to the mean and SD in the HC subjects. The relationship between both z-scores was tested by linear regression (including constant term) of the PET-PSPRP expression z-score with the MRI-SVM-PSP score as independent variable.

Receiver operating characteristic (ROC) analysis was used to characterise the utility for PSP detection separately for the MRI-SVM-PSP z-score and the PET-PSPRP expression z-score. DeLong’s test was used to compare the areas under the ROC curves.³³ ROC analyses were first performed including all PSP patients. Additional analyses were performed including only PSP-RS, only vPSP (all subtypes), only cortical vPSP or only subcortical vPSP.

Cut-off values for dichotomisation of the z-scores were obtained from the ROC curves according to Youden’s criterion. Contingency tables were used to characterise balanced accuracy,

sensitivity and specificity, separately for both z-scores. McNemar’s method was applied to the 2×2 contingency table of the predictions (PSP or HC) by the dichotomised MRI-SVM-PSP z-score versus the dichotomised PET-PSPRP expression z-score to test their difference in the proportion of PSP predictions for significance.

To examine whether the combination of the MRI-SVM-PSP z-score with the PET-PSPRP expression z-score could enhance PSP detection, their sum (MRI-SVM-PSP z-score+PET-PSPRP expression z-score) and product (MRI-SVM-PSP z-score×PET-PSPRP expression z-score) as well as a binary logistic regression model of the group label (PSP or HC) with the two z-scores as predictors were tested. The same ROC analyses were applied as for each z-score alone.

Additionally, decision tree analyses were used, employing ‘classification and regression trees’ (CART). A major feature of CART is that each tree branching results in two subgroups, simplifying the interpretation compared with other decision tree variants that allow more than two subgroups at each branching. The growth method of CART aims at minimum intranode impurity, that is, maximum intranode homogeneity (optimally, all cases of a terminal node belong to the same category). The measure of impurity to be minimised by the CART was the Gini-Simpson diversity index, which is the standard for categorical dependent variables. The Gini-Simpson index of a node characterises the rate of false categorisation when randomly selected cases are randomly categorised according to the distribution of the categories within the node.³⁴ The minimum improvement in the Gini-Simpson index required to split a node was set to the default value of 0.0001. The primary CART analysis included all PSP patients and was free to select the predictor for the first branching (MRI-SVM-PSP or PET-PSPRP expression z-score). Secondary CART analyses were restricted to patients with PSP-RS or vPSP (all subtypes) in order to test for a potential impact of the clinical phenotype on the utility of sequential MRI and FDG-PET. The secondary CART analyses were also free regarding the selection of the predictor for the first branching. A tertiary CART analysis again included all PSP patients but was forced to select the PET-PSPRP expression z-score for the first branching in order to test for an added value of MRI if FDG-PET has already been performed. For PSP detection independent

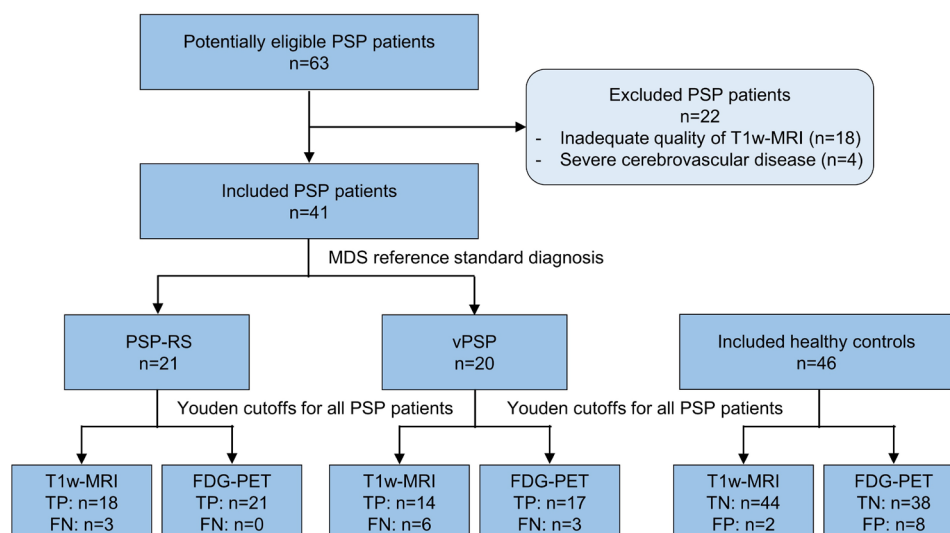


Figure 1 Flow of study participants. FDG-PET, [¹⁸F]fluorodeoxyglucose-positron emission tomography; FN, false negative; FP, false positive; PSP, progressive supranuclear palsy; RS, Richardson’s syndrome; TN, true negative; TP, true positive; vPSP, variant PSP.

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Table 1 MRI-SVM-PSP z-score and PET-PSRP expression z-score and their performance in the discrimination of the PSP patients (all, PSP-RS, vPSP, cortical vPSP and subcortical vPSP) from the HC subjects

	All PSP (n=41)	PSP-RS (n=21)	vPSP (n=20)	Cortical vPSP (n=11)	Subcortical vPSP (n=9)
z-score, mean±SD					
MRI-SVM-PSP	2.85±1.63	3.38±1.42	2.29±1.69	2.19±1.87	2.42±1.50
PET-PSRP expression	2.14±1.51	2.57±1.23	1.69±1.67	1.68±1.83	1.71±1.56
Area under ROC curve (95% CI)					
MRI-SVM-PSP	0.910 (0.848 to 0.973)	0.950 (0.891 to 1.000)	0.868 (0.764 to 0.972)	0.832 (0.665 to 0.999)	0.913 (0.819 to 1.000)
PET-PSRP	0.894 (0.826 to 0.963)	0.952 (0.908 to 0.996)	0.834 (0.726 to 0.942)	0.812 (0.662 to 0.963)	0.860 (0.738 to 0.981)
DeLong P value	0.71	0.95	0.63	0.85	0.50
Youden cut-off for dichotomisation of z-scores					
MRI-SVM-PSP	2.013	2.327	2.013		
PET-PSRP	0.516	0.728	0.516		
McNemar's test of dichotomised z-scores (MRI-SVM-PSP vs PET-PSRP expression)					
P value	0.012	0.012	0.049	0.092	0.057
Balanced accuracy in % (95% CI)					
MRI-SVM-PSP	86.9 (79.9 to 93.8)	91.8 (84.0 to 99.5)	82.8 (72.4 to 93.3)	84.2 (70.7 to 97.7)	81.2 (65.5 to 96.9)
PET-PSRP	87.6 (80.9 to 94.4)	92.4 (87.2 to 97.6)	83.8 (74.3 to 93.3)	82.2 (69.6 to 94.8)	85.7 (74.1 to 97.4)
Sensitivity in % (95% CI)					
MRI-SVM-PSP	78.0 (63.9 to 88.8)	85.7 (67.0 to 96.2)	70.0 (48.3 to 86.8)	72.7 (43.5 to 92.4)	66.7 (34.5 to 90.5)
PET-PSRP	92.7 (82.1 to 98.1)	100 (83.9 to 100)	85.0 (65.6 to 96.0)	81.8 (53.7 to 96.7)	88.9 (59.5 to 99.3)
Specificity in % (95% CI)					
MRI-SVM-PSP	95.7 (87.2 to 99.3)	97.8 (90.8 to 99.9)	95.7 (87.2 to 99.3)	95.7 (87.2 to 99.3)	95.7 (87.2 to 99.3)
PET-PSRP	82.6 (70.0 to 91.6)	84.8 (72.6 to 93.2)	82.6 (70.0 to 91.6)	82.6 (70.0 to 91.6)	82.6 (70.0 to 91.6)

Among the 11 patients with cortical vPSP, 10 presented with predominant corticobasal syndrome and 1 with predominant frontal presentation. From the nine patients with subcortical vPSP, eight presented with predominant parkinsonism and one with progressive gait freezing. The Youden cut-offs optimised for the detection of vPSP (independent of the vPSP subtype) were also applied for the detection of cortical vPSP and subcortical vPSP separately. The small sample size of the latter subgroups did not allow reliable determination of specific cut-offs.

HC, healthy controls; PET, positron emission tomography; PSRP, progressive supranuclear palsy-related pattern; RS, Richardson's syndrome; SVM, support vector machine; vPSP, variant PSP.

of phenotype (primary and tertiary CART), the minimum size of parent and child nodes was set to 30 and 15, respectively, to account for the rather small sample size. For detection of PSP-RS or vPSP (secondary CART), with consideration for a further reduction of the sample size, the minimum sizes were set at 20 and 10. The maximum depth was set to two for all CART.

IBM SPSS (V.27) was used for the statistical analyses.

RESULTS

The flow of study participants is shown in [figure 1](#).

Mean z-scores in PSP patients are given in [table 1](#). All differences compared with the HC subjects (z-score=0.0±1.0 by definition) were significant (all p<0.001).

The PET-PSRP expression z-score was significantly correlated with the MRI-SVM z-score (standardised regression coefficient $\beta=0.660$, p<0.001, [figure 2A](#)). However, the regression model explained only 43% of its variance, leaving room for improvement by the combination of both z-scores.

ROC results are given in [table 1](#) and [figure 2B](#). The area under the ROC curve was about 0.90 for detection of PSP independent of the phenotype, about 0.95 for detection of PSP-RS and about 0.85 for detection of vPSP, without significant differences between MRI-SVM-PSP and PET-PSRP expression z-score (all p≥0.63, [table 1](#)). Balanced accuracy was very similar for both z-scores, about 87% for detection of PSP independent of the subtype, approximately 92% for detection of PSP-RS and about 83% for detection of vPSP ([table 1](#)). However, the MRI-SVM-PSP z-score provided about 13% higher specificity and about 15% lower sensitivity compared with the PET-PSRP expression z-score, independent of the PSP phenotype ([table 1](#)).

Binary logistic regression of the group label (PSP vs HC) included both, the MRI-SVM-PSP z-score (coefficient B=0.814, p<0.001) and the PET-PSRP expression z-score (B=1.122, p=0.004) in addition to a constant term (B=-2.022, p<0.001). The regression model resulted in a slightly higher area under the ROC curve of 0.934 (95% CI 0.883 to 0.984) for the detection of PSP in the whole sample compared with each z-score alone, but the differences did not reach significance (p≥0.15). Balanced accuracy, sensitivity and specificity of the regression model were 86.0%, 82.9% and 89.1%, respectively. The sum of the two z-scores achieved an area under the ROC curve of 0.931 (95% CI 0.878 to 0.983), their product achieved 0.835 (95% CI 0.736 to 0.934).

Primary and secondary CART selected the MRI-SVM-PSP z-score for the first branching and the PET-PSRP expression z-score for a second split of the subgroup with normal MRI-SVM-PSP z-score after the first split, independent of the phenotype ([figure 3A](#), [figure 4](#)). In the whole sample, the primary CART achieved balanced accuracy, sensitivity and specificity of 89.0%, 97.6% and 80.4%, respectively. When the analysis was restricted to PSP-RS/vPSP (secondary CART), balanced accuracy, sensitivity and specificity were 91.3/87.7%, 100/95.0% and 82.6/80.4%, respectively. The tertiary CART is shown in [figure 3B](#).

DISCUSSION

The primary finding of this study was that the MRI-SVM-PSP score and the PET-PSRP expression score provide about the same overall (balanced) accuracy for the detection of PSP, but with a relative shift towards higher specificity of the MRI score and higher sensitivity of the PET score, suggesting that FDG-PET

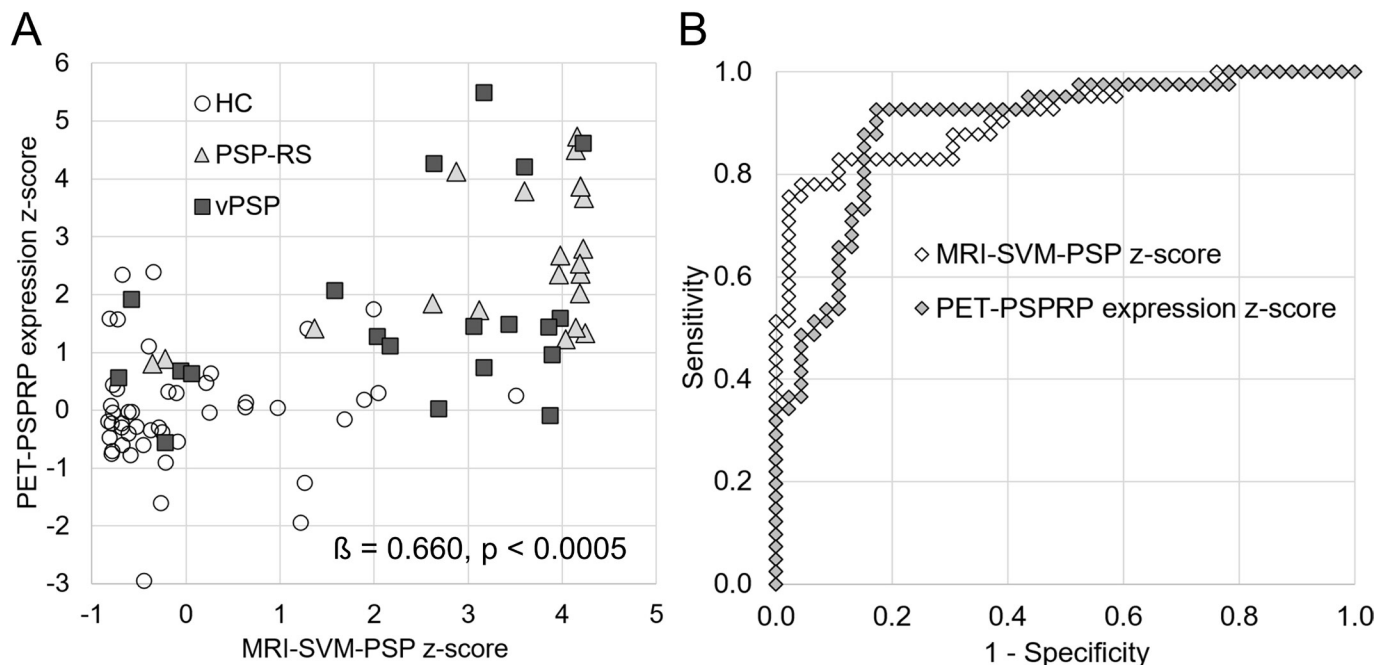


Figure 2 Scatter plot of the PET-PSRPP expression z-score versus the MRI-SVM-PSP z-score (A) and ROC curves for the detection of PSP independent of the phenotype (B). HC, healthy control; PET, positron emission tomography; PSRPP, progressive supranuclear palsy-related pattern; ROC, receiver operating characteristic; SVM, support vector machine; vPSP, variant PSP.

provides most added value in patients with normal (or inconclusive) MRI. This was confirmed by the primary decision tree analysis, which selected the MRI-SVM-PSP z-score for stratification at the first branching, followed by the PET-PSRPP expression z-score to identify PSP patients among those categorised

as normal at the MRI-based split. The PET-PSRPP expression z-score identified 9 of 10 (90%) of the PSP patients with normal MRI (figure 3A). Only a single of the 41 PSP patients was misclassified as normal by the decision tree, resulting in 97.6% sensitivity.

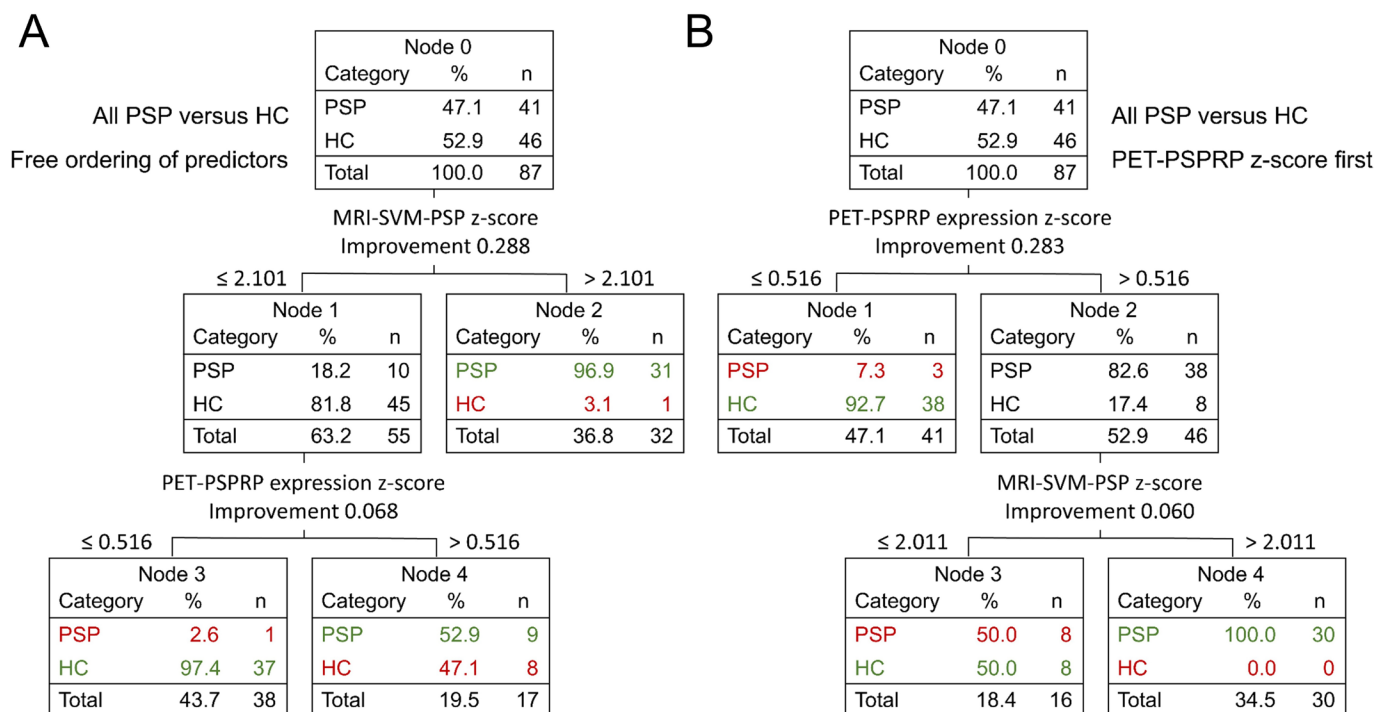


Figure 3 Decision tree analysis to combine the MRI-SVM-PSP z-score and the PET-PSRPP expression z-score for the detection of PSP independent of the phenotype. The primary decision tree (which was free to select a predictor for the first branching) selected the MRI-SVM-PSP z-score (A). (B) The tertiary decision tree that was forced to select the PET-PSRPP expression score for the first branching. Correctly classified cases are shown in green, falsely classified cases are shown in red. HC, healthy controls; PET, positron emission tomography; PSRPP, progressive supranuclear palsy-related pattern; SVM, support vector machine.

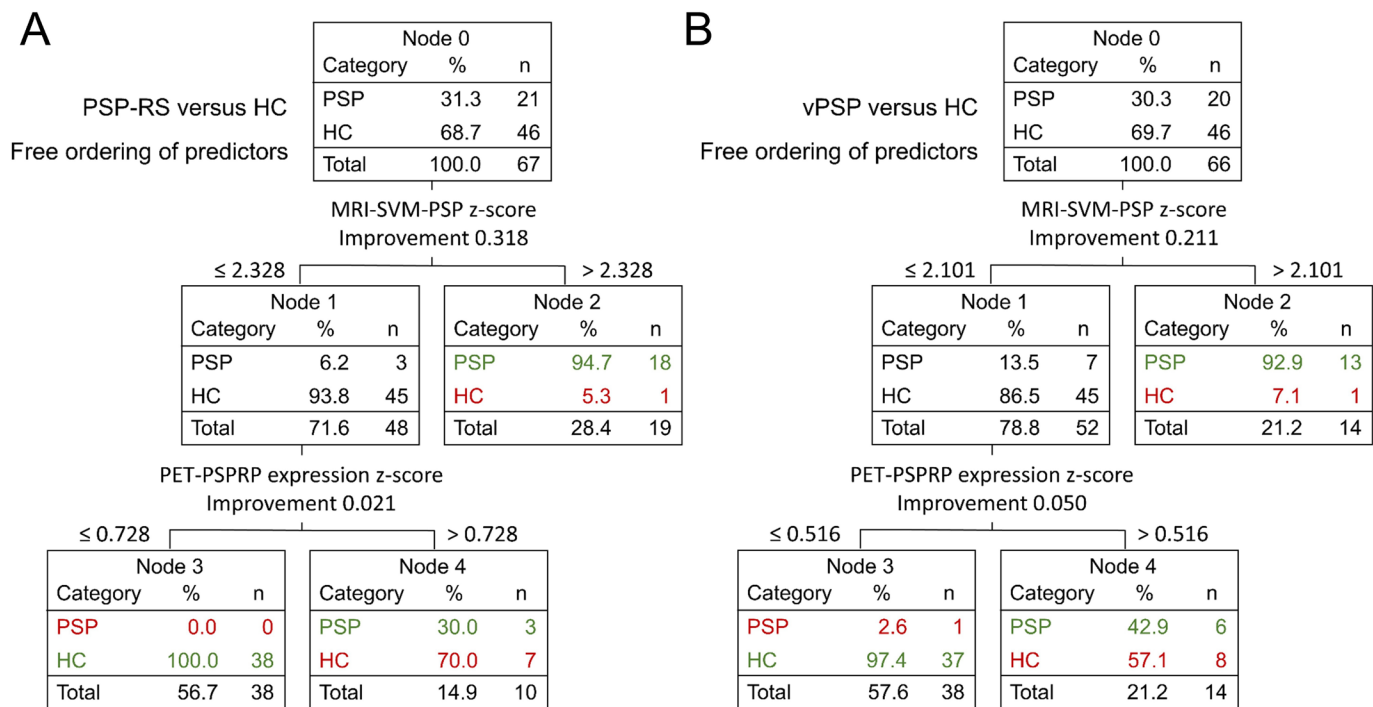


Figure 4 Secondary decision tree analyses to combine the MRI-SVM-PSP z-score and the PET-PSPRP expression z-score for the detection of PSP-RS (A) or vPSP (B). Correctly classified cases are shown in green, falsely classified cases are shown in red. HC, healthy controls; PET, positron emission tomography; PSPRP, progressive supranuclear palsy-related pattern; RS, Richardson's syndrome; SVM, support vector machine; vPSP, variant PSP.

Both z-scores were provided as predictors to the primary decision tree on equal footing, that is, without any bias to prefer one or the other for the first branching. The fact that the decision tree preferred the MRI-SVM-PSP z-score for the first branching makes the model compatible with clinical practice. Structural MRI is part of the basic diagnostics in patients suspected to have PSP to rule out secondary causes¹⁶ and, therefore, always should be performed prior to FDG-PET. Furthermore, procedure guideline for brain FDG-PET recommend that the findings of structural brain imaging, preferably MRI, are taken into account when interpreting brain FDG-PET in order to avoid misinterpretation of structural/vascular lesions (and their at-distance effects on the cerebral glucose metabolism) as indication of a primary neurodegenerative disease.³⁵ Thus, the primary decision tree model obtained in this study is ready for use in clinical practice.

The finding of about the same (balanced) accuracy with higher specificity of the MRI-SVM-PSP z-score and higher sensitivity of the PET-PSPRP expression z-score was independent of the PSP phenotype (table 1). Furthermore, the secondary decision trees for the detection of PSP-RS or vPSP also selected the MRI-SVM-PSP z-score for the first branching (figure 4). The improvement of within-node homogeneity (Gini-Simpson diversity index) by the MRI-SVM-PSP z-score at the first branching was somewhat smaller for the detection of vPSP compared with PSP-RS (0.211 vs 0.318), and the improvement by the PET-PSPRP expression z-score at the second branching was somewhat larger for the detection of vPSP (0.050 vs 0.021). This suggests that the added value of FDG-PET in patients with normal (or inconclusive) MRI is somewhat larger in vPSP than in PSP-RS.

Higher sensitivity of FDG-PET for PSP detection compared with structural MRI might be explained by neuronal dysfunction preceding brain volume loss resulting in greater and more consistent changes in FDG-PET compared with MRI-based volumetry at early PSP stages.³⁶ Concerning lower specificity of FDG-PET, retrospective inspection identified frontal widening of the longitudinal fissure as a major source of false positive FDG-PET.¹⁰ Differences in added

value of FDG-PET between vPSP and PSP-RS might be explained by phenotype-specific spatiotemporal patterns in FDG-PET and MRI-based volumetry.^{37 38}

The head-to-head comparison of MRI-based and FDG-PET-based PSP detection in the current study was restricted to SVM classification of individual atrophy profiles from fully automatic MRI volumetry and fully automatic covariance pattern expression analysis of FDG-PET. However, better specificity of MRI and better sensitivity of FDG-PET are also true when comparing visual interpretation of the two modalities. In the same subjects as included in the current study, visual reading of MRI based on descriptive signs (hummingbird, morning glory, Mickey-Mouse) and supported by automatic volumetry (same compartments as used as input to the SVM) achieved balanced accuracy, sensitivity and specificity of 74.9%, 58.5% and 91.3% for the detection of PSP independent of the phenotype (majority vote of 3 readers).¹⁸ Visual interpretation of FDG-PET supported by univariate voxel-based testing achieved balanced accuracy, sensitivity and specificity of 72.9%, 74.0% and 71.7%, respectively (majority vote of 5 readers).¹⁰ Thus, the primary conclusion of this study, that is, that FDG-PET is most beneficial in patients with normal or inconclusive MRI, will also apply if the fully automatic multivariate methods are not available so that the image interpretation is based on visual inspection (without or with support by univariate quantitative/statistical features). Taken together, these findings indicate that the main conclusion applies regardless of the phenotypic range present in a specific setting and regardless of the approach utilised to assess PSP-typical patterns in T1w-MRI and FDG-PET. However, future guidelines should recommend the use of automatic multivariate techniques to support the interpretation of T1w-MRI and FDG-PET, as these techniques are clearly superior to visual inspection regarding the detection (and exclusion) of PSP features.

A literature search (online supplemental material) identified only two studies reporting results that allow comparison of T1w-MRI and FDG-PET regarding PSP detection.^{39 40} Park *et*

*al*³⁹ found similar sensitivity of T1w-MRI and FDG-PET in 14 patients with probable PSP according to the NINDS-SPSP clinical criteria³²: both, midbrain atrophy in T1w-MRI and midbrain hypometabolism in FDG-PET were detected in 12 patients (86%).³⁹ Zhao *et al*⁴⁰ found enlargement of the third ventricle in 15 (79%) and midbrain atrophy in 13 (68%) of 19 patients with MDS PSP diagnosis. Among the 13 patients who also underwent brain FDG-PET, 12 (92%) showed midbrain hypometabolism.⁴⁰ The authors concluded that FDG-PET provides better sensitivity for the detection of PSP, particularly at early clinical stages of the disease.⁴⁰ The analysis of T1w-MRI in this previous study was restricted to the visual detection of PSP-typical signs (hummingbird) and to planimetric measures. The analysis of FDG-PET was conducted using voxel-based statistical testing against an FDG-PET normal database. Multivariate techniques for the automatic classification of T1w-MRI and/or FDG-PET were not tested nor was any attempt made to combine MRI and PET or to identify a potential added value of FDG-PET after MRI or vice versa.

The current study has limitations. First, the patient sample was rather small. To some extent, this is explained by the fact that almost a third of the eligible PSP patients had to be excluded because the T1w-MRI was not adequate for morphometric/volumetric analyses, most often due to too large spacing between slices.¹⁸ Future guidelines on the diagnosis of parkinsonian syndromes should recommend that T1w-MRI is performed with sufficient quality to allow reliable morphometric/volumetric analyses. Second, the rather small sample size did not allow splitting the dataset into training and test data. Thus, the cut-offs on the z-scores were determined in the same sample to which they were applied. This might have resulted in overly optimistic performance estimates due to overfitting. However, this applies to both z-scores and, therefore, most likely did not affect the findings with respect to their relative performance or with respect to their combination. Third, there was considerable variability of the time interval between PET and MRI in the PSP patients (due to the retrospective nature of the study). However, the PSP sample was rather balanced with respect to the sequence of PET and MRI so that a relevant bias to favour one modality over the other caused by their order of succession seems unlikely.

In conclusion, the current study indicates a clinically useful added value of brain FDG-PET beyond T1w-MRI for the diagnosis of PSP in patients with normal or inconclusive T1w-MRI, irrespective of the clinical presence of a PSP-RS or a vPSP predominance type, and irrespective of the methods used to test T1w-MRI and FDG-PET for PSP-typical features. The added value by FDG-PET might be somewhat larger in vPSP than in PSP-RS. If confirmed in independent patient samples, these findings might guide decision-making on sequential MRI and FDG-PET imaging in clinically unclear patients with suspected PSP.

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Correction notice In August 2024, this paper was resupplied as open access.

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Contributors RB: design, execution, analysis, drafting of the manuscript. H-JH, FW, GB, MB, IA, CB, MK, JL, NM, SS, OS, HB, MPW and GH: design, execution, analysis, editing of final version of the manuscript. MP-N, AD, SK, AR, AQ, SR, J-JR, CS, PGS and DE: execution, analysis, editing of final version of the manuscript.

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Competing interests H-JH has used atlas-based volumetric MRI analysis in industry-sponsored research projects. CB received a grant from the Hilde-Ulrichs-Stiftung, served as a consultant for Bial, Hormosan Pharma, Merz Pharmaceuticals and Zambon and received honoraria for scientific presentations from Abbvie, Bial, Stada Pharma, TAD Pharma, UCB Pharma and Zambon. MP-N received lecture fees from Abbott, Abbvie, Boston Scientific and served as consultant for Medtronic, Boston Scientific, Abbott, Zambon and Abbvie. SK was funded by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) under Germany's Excellence Strategy within the framework of the Munich Cluster for Systems Neurology (EXC 2145 SyNergy-ID 390857198), the Ehrmann Foundation and the Lüneburg Heritage. SK receives research funding from CurePSP and reports travel support from Life Molecular Imaging outside the submitted work. MK received honoraria for scientific presentations from Abbvie and Ever Pharma. JL reports speaker fees from Bayer Vital, Biogen, Eisai, TEVA and Roche, consulting fees from Axon Neuroscience and Biogen, author fees from Thieme medical publishers and W. Kohlhammer medical publishers and is inventor in a patent 'Oral Phenylbutyrate for Treatment of Human 4-Repeat Tauopathies' (EP 23 156 122.6) filed by LMU Munich. In addition, he reports compensation for serving as chief medical officer for MODAG, is beneficiary of the phantom share program of MODAG and is inventor in a patent 'Pharmaceutical Composition and Methods of Use' (EP 22 159 408.8) filed by MODAG, all activities outside the submitted work. J-JR received speaker honoraria from GE Healthcare. OS received research support from Life Molecular Imaging. HB received reader honoraria from Life Molecular Imaging and speaker honoraria from Novartis/AAA. MPW received speaker or consultancy honoraria from

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Supplementary Material

Added value of FDG-PET for detection of progressive supranuclear palsy in patients with normal or inconclusive MRI

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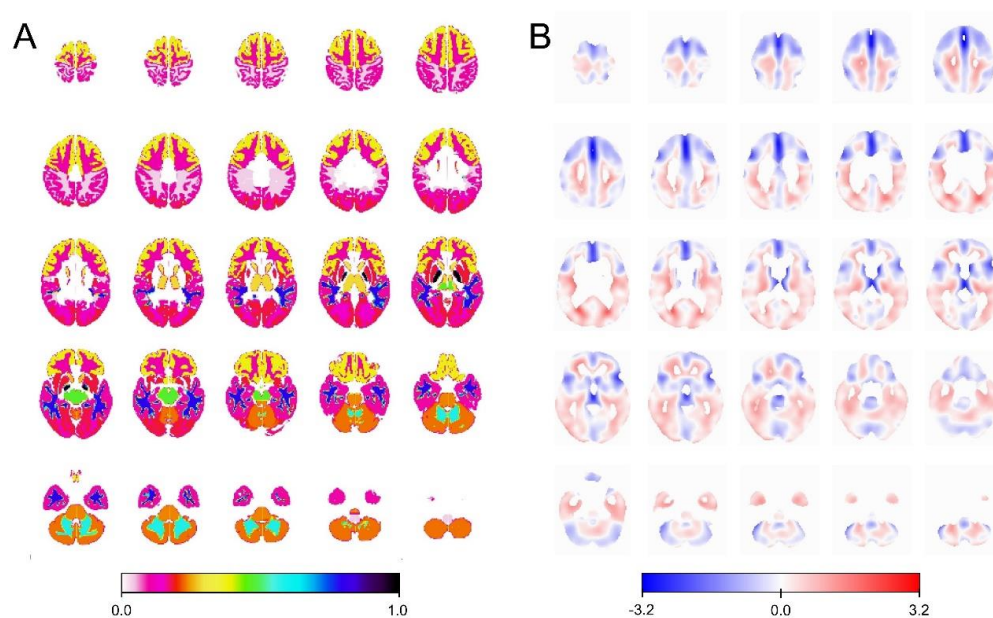
Literature search

A literature search on PubMed linking (AND) the search terms for MRI and PSP given below with a standard search strategy for FDG-PET¹ identified a total of 43 publications. From these, 41 publications were not relevant regarding the comparison of T1w-MRI and FDG-PET for the detection of PSP in individual patients: case report (≤ 3 patients): $n = 13$, review: $n = 7$, no PSP patients included: $n = 4$, other than T1-weighted MRI (e.g. functional MRI, diffusion- or T2-weighted): $n = 5$, PET with other tracer than FDG: $n = 1$, incomplete assessment of FDG-PET (restricted to the presence of midbrain hypometabolism, the ‘pimple’ sign²): $n = 1$, no data on diagnostic performance metrics reported nor data that allow to compute performance metrics retrospectively (e.g., voxel-based group level analyses to compare the atrophy pattern from voxel-based morphometry with the hypometabolism pattern from voxel-based statistical testing of FDG-PET,³ or studies on brain structural and/or metabolic signatures of specific symptoms,⁴ or prediction of PET findings from MRI findings⁵): $n = 9$, or lack of independent gold standard diagnosis (i.e., imaging-based diagnosis only): $n = 1$.

Only two studies reported results that allow comparison of T1w-MRI and FDG-PET of the brain regarding the detection of PSP.^{6,7}

Search term MRI: (magnetic resonance imaging) OR MRI OR (nuclear magnetic resonance) OR NMR OR NMRI OR (magnetic resonance tomography) OR MRT

Search term PSP: (supranuclear palsy, progressive) OR PSP OR (Richardson’s syndrome) OR (Steele-Richardson-Olszewski syndrome)



Supplementary Figure 1 Compartments used as input for the SVM classification of T1w-MRI, color-coded by the SVM weights⁸ (A), and PSPRP used for the classification of FDG-PET⁹ (B).

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Supplementary Material

Added value of FDG-PET for detection of progressive supranuclear palsy in patients with normal or inconclusive MRI

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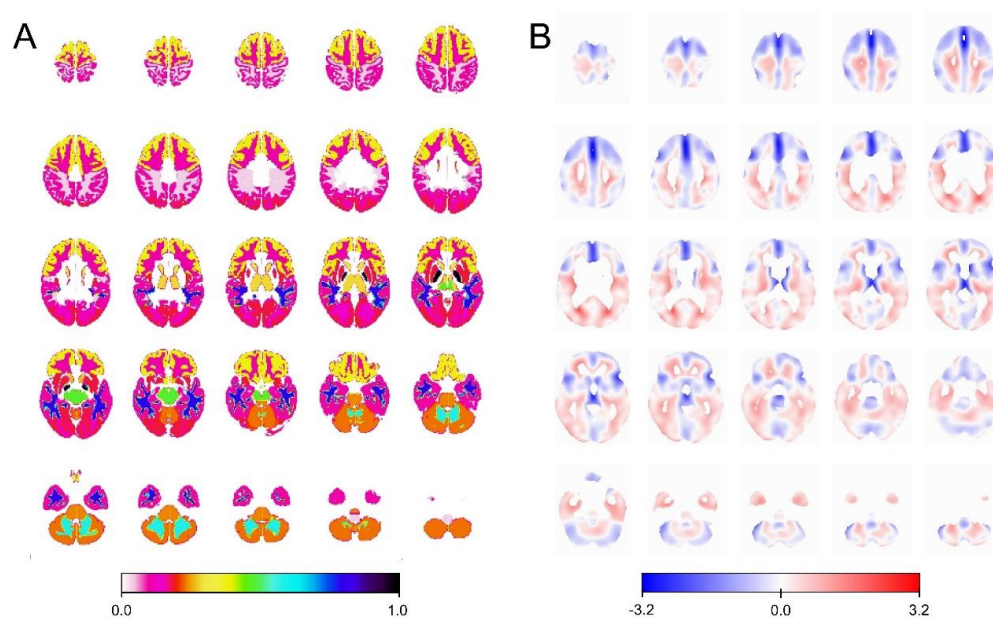
Literature search

A literature search on PubMed linking (AND) the search terms for MRI and PSP given below with a standard search strategy for FDG-PET¹ identified a total of 43 publications. From these, 41 publications were not relevant regarding the comparison of T1w-MRI and FDG-PET for the detection of PSP in individual patients: case report (≤ 3 patients): $n = 13$, review: $n = 7$, no PSP patients included: $n = 4$, other than T1-weighted MRI (e.g. functional MRI, diffusion- or T2-weighted): $n = 5$, PET with other tracer than FDG: $n = 1$, incomplete assessment of FDG-PET (restricted to the presence of midbrain hypometabolism, the ‘pimple’ sign²): $n = 1$, no data on diagnostic performance metrics reported nor data that allow to compute performance metrics retrospectively (e.g., voxel-based group level analyses to compare the atrophy pattern from voxel-based morphometry with the hypometabolism pattern from voxel-based statistical testing of FDG-PET,³ or studies on brain structural and/or metabolic signatures of specific symptoms,⁴ or prediction of PET findings from MRI findings⁵): $n = 9$, or lack of independent gold standard diagnosis (i.e., imaging-based diagnosis only): $n = 1$.

Only two studies reported results that allow comparison of T1w-MRI and FDG-PET of the brain regarding the detection of PSP.^{6,7}

Search term MRI: (magnetic resonance imaging) OR MRI OR (nuclear magnetic resonance) OR NMR OR NMRI OR (magnetic resonance tomography) OR MRT

Search term PSP: (supranuclear palsy, progressive) OR PSP OR (Richardson’s syndrome) OR (Steele-Richardson-Olszewski syndrome)



Supplementary Figure 1 Compartments used as input for the SVM classification of T1w-MRI, color-coded by the SVM weights⁸ (A), and PSRP used for the classification of FDG-PET⁹ (B).

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