PET Radioligands Reveal the Basis of Dementia in Parkinson’s Disease and Dementia with Lewy Bodies

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Abstract
Background: Effective therapies for dementia with Lewy bodies (DLB) and Parkinson’s disease (PD) dementia will require accurate diagnosis and an understanding of the contribution of distinct molecular pathologies to these diseases. We seek to use imaging biomarkers to improve diagnostic accuracy and to clarify the contribution of molecular species to cognitive impairment in DLB and PD. Summary: We have performed cross-sectional and prospective cohort studies in subjects with DLB, PD with normal cognition, PD with mild cognitive impairment and PD with dementia, contrasted with Alzheimer’s disease (AD) and healthy control subjects (HCS). Subjects underwent formal neurological examination, detailed neuropsychological assessments, MRI and PET scans with the radioligands alropane (a dopamine transporter, DAT) and Pittsburgh compound B (PiB; β-amyloid). Putamen DAT concentrations were similar in DLB and PD and differentiated them from HCS and AD. Decreased caudate DAT concentration related to functional impairment in DLB but not PD. PiB uptake was greatest in DLB. However, cortical PiB retention was common in PD and predicted cognitive decline. PET imaging of tau aggregates holds promise both to clarify the contribution of tau to cognitive decline in these diseases and to differentiate DLB and PD from the parkinsonian tauopathies. Key Messages: Together, DAT and amyloid PET imaging discriminate DLB from PD and from other disease groups and identify pathological processes that contribute to their course. Multimodal PET imaging has the potential to increase the diagnostic accuracy of DLB and PD in the clinic, improve cohort uniformity for clinical trials, and serve as biomarkers for targeted molecular therapies.

Dementia with Lewy bodies (DLB) is the second most common dementia after Alzheimer’s disease (AD) [1]. Its clinical features, which include dementia associated with the motor manifestations of parkinsonism, visual hallucinations and fluctuations of cognition, overlap significantly with the clinical features that arise in the context of
dementia in idiopathic Parkinson’s disease (PD) [2]. DLB and PD dementia (PDD) are differentiated by the relative timing of dementia and parkinsonism, with PDD arising in the context of well-established idiopathic PD, after at least 1 year of motor symptoms, whereas an early onset of dementia is more consistent with a diagnosis of DLB [1]. Both DLB and PDD are characterized neuropathologically by the accumulation of α-synuclein aggregates in Lewy bodies and Lewy neurites in the brain, with cortical Lewy bodies required for a neuropathological diagnosis of DLB but common in PDD. Given their clinical and neuropathological similarities, DLB, PDD and PD have been aggregated conceptually under the broad umbrella of Lewy body disease [2].

Despite the value of the DLB Consortium criteria [1], the accuracy of a clinical diagnosis of DLB remains low. Sensitivity of the DLB consensus criteria has been found to be quite variable, ranging from 12 to 88% across different academic centers [3], and is greatest with inclusion of supportive diagnostic features, including PET imaging demonstrating dopamine cell loss, neuroleptic sensitivity or REM sleep behavioral disorder. In contrast, the specificity of the criteria is higher, ranging from 79 to 100%. The repercussions of an incorrect diagnosis are significant, including risk for neuroleptic exposure, lost opportunity for treatment, and inappropriate exclusion from and inclusion in clinical trials.

Molecular imaging tools hold promise for improving the diagnostic accuracy of DLB. For example, dopamine transporter (DAT) imaging with SPECT or PET can confirm neurodegenerative parkinsonism [4]. Using [11C]al-tropane, a DAT ligand with selective affinity for the DAT over the norepinephrine transporter or the serotonin transporter (7 nmol affinity, 25-fold greater affinity for the DAT over the serotonin transporter) [5], we have begun to explore the relation of DAT levels with clinical features of DLB and PD [6].

In addition, comorbid AD pathology is common in DLB, such that approximately 85% of cases contain abundant amyloid plaques [7, 8]. Aggregates of tau in the form of neurofibrillary tangles are also observed [8]. The development of PET ligands that label brain amyloid, such as [11C]Pittsburgh compound B ([11C]PiB), has provided a noninvasive method to explore the contribution of amyloid deposition during life to the clinical features and clinical course of DLB and PD [9]. Amyloid imaging reliably labels fibrillar forms of β-amyloid, including cored plaques, some diffuse plaques and amyloid angiopathy. As negative scans are exceedingly rare in AD, this has proven to be a valuable molecular imaging tool. The clinical relevance of concomitant AD pathology in DLB remains a topic of active investigation. Some studies have found that amyloid deposition impacts the clinical phenotype [10]. In addition, we and other groups have found that amyloid deposition hastens cognitive decline in PD [11, 12].

Recently, several PET ligands have been developed to image paired helical filaments of tau, which accumulate in neurofibrillary tangles in AD. One such agent, [18F]T807, has a very high affinity for paired helical filaments of tau and a selectivity for tau over β-amyloid of >27-fold [14]. The application of [18F]T807 imaging to PD and DLB provides a powerful new method to evaluate, during life, the contribution of paired helical filaments of tau to cognitive impairment and clinical phenotype. In addition, to the extent that [18F]T807 may bind tau aggregates in the parkinsonian tauopathies, progressive supranuclear palsy (PSP) and cortical basal ganglionic degeneration, [18F]T807 also has the potential to discriminate between parkinsonian diseases of synuclein and tau.

Here, we report our experience with DAT imaging and amyloid imaging in cross-sectional and longitudinal studies of DLB and PD, as well as our early experience with [18F]T807 PET imaging.

**Methods**

**Subjects**

Subjects were recruited from the Massachusetts General Hospital Movement Disorders and Memory Disorders Units and gave informed consent to participate in this research study according to the protocol approved by the Partners Health Care Inc. Institutional Review Board. PD subjects fulfilled criteria for a diagnosis of idiopathic PD according to the UK Parkinson Disease Society Brain Bank Research Center’s clinical diagnostic criteria [15]. Criteria for PD with mild cognitive impairment (PD-MCI) were consistent with those of the Movement Disorder Society [16], requiring a cognitive complaint, such as impairment of short-term memory, plus at least one cognitive test score that fell in the impaired range. We used the Movement Disorder Society criteria for PDD [17] and the DLB Consortium consensus criteria for DLB [1]. Each subject underwent standardized neurological examination, cognitive testing and PET imaging.

**Clinical Evaluation**

Functional status was assessed by the Clinical Dementia Rating Scale Sum of Boxes (CDR-SB). Neuropsychological testing was also acquired, as described in Gomperts et al. [11]. Motor function was assessed with the Hoehn and Yahr stage and the motor subscale of the Unified Parkinson Disease Rating Scale.

For DAT imaging, we enrolled 81 subjects, including 29 with PD without dementia (20 cognitively normal, 9 PD-MCI), 12...
DLB, 10 PDD and 6 AD. Twenty-four subjects served as a healthy control subject (HCS) group. Of this cohort, 19 PD, 10 DLB and 17 HCS also underwent structural brain MRI for Freesurfer-derived partial volume correction of PET data. Detailed altropane PET methods are described in Marquie et al. [6].

For PiB imaging, we studied a total of 74 subjects, including 29 PD with normal cognition (PD-nl), 14 PD-MCI, 12 PDD and 18 DLB. Of these, 20 PD-nl, 5 PD-MCI, 10 PDD and 9 DLB had also been imaged with altropane. Data from these subjects were compared with those of a separately collected cohort of 85 HCS. Detailed cross-sectional PiB PET methods are described in Gomperts et al. [18].

All HCS subjects had a normal neurological examination, a CDR-SB score of 0 and normal cognition, with the Blessed Dementia Scale IMC score <3 or Mini Mental State Examination score >27.

For the prospective cohort study, 46 nondemented PD subjects underwent PiB imaging and were then followed with annual cognitive and neurological evaluations, for a mean of 3.9 ± 1.9 visits (range: 1–7 visits). Of the 46 subjects, 35 scored in normal ranges on a set of cognitive tests at study baseline, and 32 of these had no cognitive complaint. Eleven moved Movement Disorder Society criteria for MCI [16]. PD-MCI subjects had a global CDR score of at least 0.5. Eight out of 11 cases were amnestic, multidomain MCI, and 3/11 were nonamnestic, multidomain MCI. The memory box score was at least 0.5 in 10 of these subjects. The mean Mini Mental State Examination score was 29.1 ± 1.0 for the PD-nl group and 26.3 ± 2.9 for the PD-MCI group. Detailed longitudinal PiB PET methods are described in Gomperts et al. [11].

We have also acquired [18F]T807 PET imaging in a small number of PiB-imaged subjects with PD, PD-MCI, PDD, DLB, PSP and corticobasal syndrome.

**PET Imaging Acquisition and Analyses**

[11C]Altropane. Altropane PET images were acquired as described previously [6]. Briefly, 15 mCi [11C]altropane was injected as a bolus and followed with a 60-min dynamic acquisition. DAT concentration was estimated with specific binding of altropane, which was computed in regions of interest (ROIs) using the standarized uptake value ratio (SUVR) [19], a ratio of uptake in the target ROI to reference region measured between 40 and 60 min after injection. The pericalcarine (visual) cortex was selected as reference on the basis of its low DAT concentration after injection. The pericalcarine (visual) cortex was selected as target ROI to reference region measured between 40 and 60 min after injection. The pericalcarine (visual) cortex was selected as target ROI to reference region measured between 40 and 60 min after injection. The pericalcarine (visual) cortex was selected as target ROI to reference region measured between 40 and 60 min after injection.

For [18F]T807 data acquisition, administration of 10 mCi of radiotracer was followed by a 20-min acquisition, beginning at 80 min after injection. We expressed [18F]T807 data as the SUVR with cerebellar reference, as previously reported [14]. SUVR images were coregistered to a magnetization-prepared rapid acquisition gradient echo sequence to generate high-resolution anatomic data. [18F]T807 PET data were spatially normalized to the MNI template brain using SPM2.

**Data Analysis**

DAT concentration in the putamen and caudate ROIs was evaluated using a backward elimination general linear model (GLM) regressed on the initial predictor pool: diagnostic group, age at imaging, education, putamen DAT (for the caudate analyses), duration of motor symptoms, levodopa-equivalent dose and the interaction of diagnosis with each of the other predictors. The cutoff p value for removal from the model was 0.01. A covariate of caudate volume and its interaction with diagnosis were included when caudate DAT concentration was the dependent variable. Tukey post hoc tests were performed as required to follow up significant diagnostic main effects. We assessed the relation of DAT concentration to CDR-SB using GLM analyses with backward elimination (p > 0.01 for removal from the model), with a pool of predictors that initially included diagnostic group, DAT concentration in the selected ROIs, interaction of diagnosis with ROIs, caudate volume, age, education, duration of motor symptoms and levodopa-equivalent dose.

Precuneus PiB retention (DVR) was compared across the diagnostic groups (PD, PD-MCI, PDD, DLB and HCS) using a backward elimination GLM with PiB as the dependent variable and with an initial pool of predictor variables which included diagnostic group, age at imaging and interactions of diagnosis with age. Cutoff p values for removal from the model were set at 0.05. Tukey-adjusted post hoc tests were performed where required to follow up significant main effects.

For longitudinal PiB analyses, longitudinal change in the CDR-SB was assessed with mixed random/fixed coefficient longitudinal regression models. We conducted backward elimination of fixed terms, which included baseline diagnosis (PD-nl/ PD-MCI) and its interaction with time, PiB DVR and its interaction with time, gender, years of education, baseline age and baseline duration of motor symptoms. The relation of PiB DVR to time to transition to a more severe diagnosis was assessed with Cox proportional hazards regression models containing the predictors of PiB DVR, baseline diagnosis and its interaction with PiB, baseline age/duration, years of education and gender. Earliest transitions from PD-nl to PD-MCI (or PDD) and from PD-MCI to PDD were both included in the analysis. PD subjects either remained stable in their diagnosis across time (considered ‘censored’ observations in the model) or transitioned to a more severe diagnosis. The assumption of proportional hazards was checked and verified.

SAS (version 9.3) and JMP Pro software were used for analyses and graphs.
Results

DAT Imaging

Putamen DAT concentration was significantly lower in the Lewy body dementia cases (DLB and PDD) than in AD and HCS (each comparison, p < 0.0001, GLM; fig. 1a). In addition, caudate DAT concentration, assessed with Freesurfer, was also lower in DLB and in PD than in HCS (each comparison, p < 0.0003; fig 1b). Adjusting for age and for putamen DAT concentration, as a measure of severity of motor system impairment, caudate DAT concentration was significantly lower in DLB than in PD (p < 0.041; fig. 1c). These data confirm the value of DAT imaging with altropane both to differentiate DLB and PDD from AD and to distinguish DLB from PD.

We also examined the relation of caudate DAT concentration to cognitive function in DLB. Adjusting for age, duration of motor symptoms and putamen DAT concentration, higher CDR-SB was associated with greater reduction in DAT concentration in DLB (R^2 = 0.84, p < 0.0001 for model). Note that a similar decline in caudate DAT concentration in non-demented PD was not associated with a similar impairment on the CDR-SB (p = 0.0008 for interaction between DAT concentration and diagnosis).
reduction in DAT concentration ($R^2 = 0.84$, $p < 0.0001$ for model; fig. 1d). In contrast, a similar reduction of caudate DAT concentration in PD was not associated with impairment in the CDR-SB ($p = 0.0008$ for interaction between DAT concentration and diagnosis). Thus, progressive dopamine denervation of the caudate appears to contribute to cognitive impairment in DLB but is not sufficient for cognitive impairment in PD.

**Amyloid Imaging**

Cortical amyloid burden measured with $[11 \text{C}]$PiB was higher in DLB than in HCS, PD or PDD ($p < 0.0001$ for model, $p < 0.002$ for each group contrast). Even so, specific cortical PiB retention was present in many PD-nl, PD-MCI and PDD subjects, as well as in approximately 30% of HCS.

We have continued to follow the PD cohort with annual neuropsychological evaluation and neurological examinations, in order to relate amyloid burden to the course of cognitive impairment in PD and DLB. As we first described in 2013 [11] and have now extended, higher PiB uptake predicted a faster conversion of diagnosis from PD-nl to PD-MCI/PDD ($n = 10/35$) and PD-MCI to PDD ($n = 5/11$; $p = 0.008$; Cox regression; fig. 2). This corresponds to a hazard of conversion of 15.0 per unit PiB (DVR). In addition, higher PiB retention significantly related to a worsening CDR-SB score for subjects with PD-MCI or PDD ($p = 0.0179$; mixed random/fixed longitudinal model). These data confirm that amyloid burden is a risk factor for dementia in PD.

**$[18 \text{F}]$T807 PET Imaging**

We have begun $[18 \text{F}]$T807 PET acquisition in subjects clinically diagnosed with PD, PDD, DLB and the parkinsonian tauopathies PSP and cortical basal ganglionic degeneration. In our preliminary experience in nondemented PD cases, $[18 \text{F}]$T807 uptake is increased in subjects with MCI. In addition, in PSP, the distribution of $[18 \text{F}]$T807 appears to correspond to the pattern of pathology, with high levels of $[18 \text{F}]$T807 binding in the globus pallidus and other affected deep gray structures. Greater sample sizes will be required to confirm these preliminary observations and determine whether $[18 \text{F}]$T807 binding relates to amyloid burden and to cognitive impairment in PD and differentiates the synucleinopathies from the parkinsonian tauopathies.

**Discussion**

The use of PET imaging to evaluate molecular neuropathologies in living patients with PD and DLB continues to provide new insights into these disorders. Consistent with prior reports with other DAT imaging ligands,
we find that reduction of putamen DAT concentration in DLB, measured with altropane, appears to provide a useful distinction from AD, where putamen DAT is relatively preserved. Furthermore, the relative loss of caudate DAT provides a means to differentiate DLB from PD. The observation that loss of caudate DAT concentration, after adjusting for putamen DAT concentration, is associated with cognitive impairment in DLB but not in PD suggests that additional pathological processes, such as α-synuclein aggregation and the presence of copathologies, make DLB brains susceptible to loss of caudate dopamine.

One such copathology, deposition of β-amyloid, is well evaluated with PiB PET imaging. In our experience and that of other studies [9], cortical amyloid deposition is common and is often high in DLB, while cortical amyloid burden is common but often lower in PDD and PD. Nonetheless, cortical amyloid deposits are associated with a significant risk for cognitive decline in PD, in a dose-dependent manner. Thus, amyloid copathology is common and deleterious in both DLB and PD. These data suggest that treatments under development for AD that target β-amyloid 42 are likely to have clinical value in DLB and PD as well.

As yet, our preliminary experience with [{18}F]T807 supports its potential as a marker of cognitive impairment in PD and as an imaging method to differentiate PD and DLB from PSP and cortical basal ganglionic degeneration. We continue to evaluate these hypotheses.

In conclusion, DAT and amyloid imaging offer a powerful approach to diagnose DLB and PDD and to explore the molecular basis of dementia in these disorders. [{18}F] T807 may prove to be a useful additional imaging modality for these purposes.

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Disclosure Statement

Dr. Gomperts has nothing to disclose.

References


