ABSTRACT

BACKGROUND AND PURPOSE: Postmortem studies of advanced PD have revealed disease-related pathology in the thalamus with an apparent predilection for specific thalamic nuclei. In the present study, we used DTI to investigate in vivo the microstructural integrity of 6 thalamic regions in de novo patients with PD relative to healthy controls.

MATERIALS AND METHODS: Forty subjects (20 with early stage untreated PD and 20 age- and sex-matched controls) were studied with a high-resolution DTI protocol at 3T to investigate the integrity of thalamic nuclei projection fibers. Two blinded, independent raters drew ROIs in the following 6 thalamic regions: AN, VA, VL, DM, VPL/VPM, and PU. FA values were then calculated from the projection fibers in each region.

RESULTS: FA values were reduced significantly in the fibers projecting from the AN, VA, and DM, but not the VPL/VPM and PU, in the PD group compared with the control group. In addition, there was a reduction in FA values that approached significance in the VL of patients with PD. These findings were consistent across both raters.

CONCLUSIONS: The present study provides preliminary in vivo evidence of thalamic projection fiber degeneration in de novo PD and sheds light on the extent of disrupted thalamic circuitry as a result of the disease itself.

ABBREVIATIONS: AN = anterior nucleus; DM = dorsomedial nucleus; FA = fractional anisotropy; PD = Parkinson disease; PU = pulvinar; SN = substantia nigra; VA = ventral anterior nucleus; VL = ventral lateral nucleus; VPL = ventral posterior lateral nucleus; VPM = ventral posterior medial nucleus

PD is a progressive neurodegenerative disorder with a slow, predictable course through vulnerable regions of subcortical and cortical gray matter. The pathologic hallmark of PD is the selective and substantial loss of dopaminergic neurons in the SN pars compacta. It is generally accepted that a reduction of dopamine in the SN pars compacta underlies dysfunction within the direct basal ganglia pathway, which results in reduced excitation of cortical neurons by the thalamus. This increased inhibition of cortical-subcortical circuits in PD results in bradykinesia and various degrees of cognitive impairment. Thus, while dysfunction of subcortical structures such as the basal ganglia and thalamus have historically been conceptualized as a consequence of SN pars compacta degeneration, there is also evidence to suggest that the thalamus may also undergo structural changes, display functional abnormalities, and be a site of direct disease pathology in PD.

While conventional MR imaging has limited utility as an investigative and diagnostic tool in PD, recent advances in neuroimaging techniques have allowed for the noninvasive in vivo assessment of PD-related pathology. For example, DTI measures within the SN have shown excellent sensitivity and specificity in differentiating de novo patients with PD from healthy controls. Recent studies support the reduction in FA values within the SN of patients with PD and also suggest that microstructural integrity of the thalamus may be altered in PD.
crostructural changes occur in areas beyond the SN, including the thalamus.16 As a technique, DTI is based upon the diffusivity of water molecules, which is variably restricted by different tissues. FA values, which are calculated from the diffusion tensor images, range from 0 to 1, with 0 representing isotropic diffusion and 1 anisotropic diffusion. In WM, where the tissue is highly organized, the free diffusion of water is limited, resulting in FA values near 1. FA values calculated in WM reflect a combination of several factors, including axonal attenuation, degree of myelination, and orientational coherence.18,19 The direction of diffusion is less restricted in gray matter, resulting in FA values near 0. It is unclear exactly what FA values represent in gray matter structures, but one hypothesis is that they reflect neuronal attenuation, with lower FA values representing a decrease in attenuation.20

Postmortem studies demonstrate that PD differentially affects specific thalamic nuclei,10,11,21-23 indicating that there could be degeneration or disruption of the microstructural integrity of the thalamus. To date, most studies utilizing DTI to investigate the thalamus have focused on the thalamus as a single entity and have not evaluated the integrity of specific nuclei or the critical cortical-subcortical fibers that project from the thalamus.15,16,24,25 Further, these studies examined relatively advanced patients with PD who had been taking antiparkinsonian medication,15,16,24,25 thus making it unclear whether differences between the patients and controls were caused by the disease, medication, or a combination of both.

In the current study, we conducted a detailed examination of the projection fibers from specific thalamic nuclei in de novo patients with PD, allowing investigation of the disease without concern for the potential effects of antiparkinsonian medication. Using a high-resolution DTI protocol at 3T, we placed small ROIs in 6 thalamic regions and used the voxels within the ROIs as seeds for fiber tracking. Similar methods have been applied previously with high intrarater reliability in a study of traumatic brain injury.26 Our hypothesis was that there would be degeneration of thalamic projection fibers in early stage, untreated PD, and that fibers from specific nuclei may be preferentially affected.

**MATERIALS AND METHODS**

**Subjects**

Forty individuals participated in this study: 20 patients with de novo PD (10 men, 10 women) and 20 age- and sex-matched healthy volunteers. There was no significant difference in age between the groups (PD = 57.9 ± 8.9 years versus control = 58.2 ± 9.6 years; P = .7). All patients were diagnosed with PD by a movement-disorders neurologist at Rush University Medical Center and met the United Kingdom PD Society Brain Bank diagnostic criteria.27,28 For the PD group, the mean (SD) Unified Parkinson Disease Rating Scale Part III motor score was 16.7 (7.6), indicating mild disease, and the mean (SD) disease duration in months was 12.8 (10.7). None of the controls reported a history of neuropsychiatric or neurologic problems. All subjects provided written informed consent for the procedures in this study, which were approved by the institutional review boards at Rush University Medical Center and the University of Illinois at Chicago.

**Image Acquisition**

To reliably perform FA analysis and fiber tracking in the thalamus, we used a high-resolution DTI protocol that was designed to minimize eddy-current-induced distortion.29 All images were collected on a 3T Signa HDx (GE Healthcare, Milwaukee, Wisconsin) using an 8-channel phased-array head coil, together with parallel imaging. The DTI data acquisition parameters were as follows: TR = 4,500 ms; TE = 82 ms; flip angle = 90°; b values = 0, 1000 seconds/mm²; diffusion directions = 27; FOV = 200 mm²; image matrix size = 256 × 256; number of sections = 15; section thickness = 4 mm; section gap = 1 mm; NEX = 4; and acceleration factor = 2. Thus, the voxel size was 0.78 × 0.78 × 4 mm. The images were collected in the axial orientation, with the top section placed approximately 4 mm above the corpus callosum. The total acquisition time of the DTI sequence was 8 minutes, 33 seconds.

To visualize the thalamus and differentiate it from surrounding structures, a set of T2-weighted images was acquired using a fast spin-echo pulse sequence in the axial plane (TR = 5000 ms; TE = 97 ms; flip angle = 90°; FOV = 200 mm²; image matrix size = 512 × 512; number of sections = 15; section thickness = 4 mm; section gap = 1 mm; NEX = 2).

To help visualize the DM, we collected axial images using a T1-weighted 3D inversion recovery fast spoiled gradient recalled pulse sequence (TR = 25 ms; TE = 3 ms; flip angle = 40°; FOV = 240 mm²; image matrix size = 256 × 256; number of sections = 120; section thickness = 1.5 mm; section gap = 0 mm; NEX = 1).

**Image Analysis**

DTIStudio (Johns Hopkins University, Baltimore, Maryland)30 was used to reconstruct the images and calculate FA values. For all subjects, each of the 15 diffusion tensor images was examined visually for artifacts related to eddy currents and motion, namely, spatial blurring, scaling, and shearing. Head motion was further assessed using AFNI31 software (http://afni.nimh.nih.gov/afni) and required to be within 1 mm. A background noise level of 125 (MR units) was applied before calculation of the pixel-wise FA maps.

**Fiber Tracking**

Fiber tracking in DTIStudio32 was used to assess projection fibers from the 6 thalamic regions. After the calculation of pixel-wise FA and eigenvector maps, fiber tracking parameters were set to exclude tracking fibers with FA values <0.20 or a turning angle >70°. For each subject, 2 circular ROIs (3 mm in diameter) were placed in the AN, VA, VL, VPL/VPM, and PU. Given its larger size, 3 ROIs of the same diameter were placed in the DM. ROIs were drawn independently by 2 raters who were blinded to patient or control status. Fig 1A shows the regional boundaries for placing the ROIs, which were based on methods described by Little and colleagues (Appendix e-1).26 VPL and VPM were considered as one region due to the difficulty visualizing their boundaries. For each thalamic region and subject, mean FA values were calculated across the entire extent of the fibers identified by the ROI seed voxels. ROIs were drawn separately for the right and left hemispheres.
Placement of ROIs in Thalamic Regions

Anterior Nucleus. The AN ROIs were defined laterally by the internal capsule, medially by the edge of the thalamus, and anteriorly by the posterior edge of the caudate head and lateral ventricle. The posterior boundary for the ROIs was defined by a horizontal line extending from the posterior edge of the anterior limb of the internal capsule. Any fiber that extended to the putamen or globus pallidus was excluded. Furthermore, any fibers that were also identified from ROIs placed in the VA, VPL/VPM, PU, or DM were excluded.

Ventral Anterior Nucleus. The anterior edges of the VA ROIs were defined by a horizontal line extending from the anterior edge of the internal capsule and extended medially the width of the internal capsule. The VA ROIs extended posterior to a horizontal line drawn from the posterior boundary of the putamen. Any fibers that were also identified by the AN ROIs were excluded.

Ventral Lateral Nucleus. The VL ROIs were defined medially by a vertical line drawn down from the medial edge of the bend of the internal capsule (the intersection between the anterior limb of the internal capsule and the body of the internal capsule), laterally by the edge of the internal capsule, anteriorly by a horizontal line extending from the posterior edge of the anterior limb of the internal capsule, and posteriorly by a horizontal line extending from the posterior boundary of the putamen. Fibers that were previously identified by either the AN or VA ROIs were excluded.

Dorsomedial Nucleus. The DM ROIs were defined medially by the edge of the thalamic body, laterally by a vertical line extending from the posterior edge of the anterior limb of the medial edge of the internal capsule, posteriorly by a horizontal line extending from the posterior edge of the posterior limb of the internal capsule, and anteriorly by a horizontal line extending from the posterior edge of the putamen. Fibers that were previously identified by the AN, VA, VL, VPL/VPM, or DM ROIs were excluded.

Ventral Posterior Lateral and Ventral Posterior Medial Nuclei. The VPL/VPM ROI was defined anteriorly by a horizontal line extending from the posterior edge of the putamen, laterally by a vertical line extending from the posterior intersection of the internal capsule and putamen, medially by a vertical line extending the intersection of the anterior limb of the internal capsule with the anterior edge of the putamen, and posteriorly by the posterior edge of the thalamus. Fibers that were also identified by the AN, VA, VL, VPL/VPM, or DM ROIs were excluded.

Pulvinar. The PU ROIs were defined medially and posteriorly by the edge of the thalamus, laterally by a vertical line extending from the medial intersection of the anterior limb of the internal capsule and body of the internal capsule, and anteriorly by a horizontal line extending from the intersection of the posterior limb of the internal capsule and putamen. Fibers that were previously identified by the AN, VA, VL, VPL/VPM, or DM ROIs were excluded.

Statistical Analysis

For each thalamic region, FA values within each group were assessed for outliers. No FA value fell more than 3 SD from its respective mean; thus, all data were included in the analyses. Next, we performed a 2-way mixed design ANOVA on the FA values for each thalamic region, with group (PD, control) as the between-subjects factor and hemisphere as the within-subjects factor (right, left). The assessment of interrater reliability for ROI placement showed strong agreement between the 2 raters, with all interclass correlation coefficients above 0.82. Because the 2-way interaction did not approach significance for any analysis, we collapsed the data across the cerebral hemispheres. For thalamic regions that differed significantly in FA values between groups, we performed a correlation analysis between FA values and Unified Parkinson Disease Rating Scale Part III scores.

RESULTS

The 2-way interaction between group and hemisphere was not significant for any of the 6 thalamic regions (all Ps > .4). The main effect of hemisphere was only significant for the VL (P = .012), with higher FA values on the left than right side. There was a significant main effect of group for 3 of the 6 thalamic regions. As presented in Fig 2, FA values were reduced in the PD group compared with the control group for fibers projecting from the AN.
Differ significantly between the groups (both the FA values in the VPL/VPM and PU projection fibers did not reduce in the PD group compared with the control group (\( P < .016 \)). VA (\( P = .009 \)) and DM (\( P = .000 \)). In addition, the reduction in FA values from the VL approached significance for the PD group compared with the control group (\( P = .065 \)). The FA values in the VPL/VPM and PU projection fibers did not differ significantly between the groups (both \( P > .4 \)).

For the 3 thalamic regions that differed significantly between the groups, we performed a correlation analysis to investigate the relationship between Unified Parkinson Disease Rating Scale Part III scores and FA values in PD. There was a significant correlation between Unified Parkinson Disease Rating Scale Part III scores and FA values in the AN (\( r = -0.487, P = .029 \)), indicating a moderate negative association; as Unified Parkinson Disease Rating Scale Part III scores (i.e., motor impairment) increased, FA values decreased. The correlations between Unified Parkinson Disease Rating Scale Part III scores and FA values in the DM, VA, and VL were not significant (all \( P > .2 \)).

**DISCUSSION**

The thalamus plays an important role in the output pathway of the basal ganglia circuitry. It not only transmits nearly all cortical inputs, including cognitive, sensory, and motor information, but also modulates and integrates this information. Further, cerebellar output influences the cortical areas involved in motor processing via the thalamus. The thalamus is organized into distinct nuclear regions, from which diffuse and specific efferent projections are sent to cortical, cerebellar, and subcortical regions.\(^3\)

The present study was the first to examine the integrity of thalamic nuclei projections in de novo PD using in vivo DTI tractography. We showed that there were reduced FA values in the fibers projecting from the AN, VA, and DM, but not the VPL/VPM or PU, in patients with PD compared with controls. Further, the reduction in FA values approached significance in the VL of the patients with PD. Although the pathologic cause of these reduced FA values is not known, it may reflect demyelination, neuronal loss, or gliosis.\(^{20,32–35}\) We also showed that there was a significant negative correlation between FA values in the AN and Unified Parkinson Disease Rating Scale Part III scores. This relationship should be investigated further in future work. Collectively, our results provide preliminary evidence of thalamic fiber degeneration in early stage PD and shed light on the extent of disrupted thalamic circuitry as a result of the disease.

Our DTI results are largely consistent with postmortem studies of the thalamus in PD. Rüb et al\(^{11}\) reported evidence of mild inclusion pathology in the AN, VA, VL, and DM of patients with mild to severe PD. Further, there was no evidence of pathology in the VPM and VPL, and relatively little pathology in the PU of these patients. However, other studies have reported that the volume and number of neurons in the AN and DM are not reduced significantly in PD.\(^{21–23}\)

The AN, VA, and DM are considered specific nuclei, in that they relay information to functionally distinct cortical regions. The AN is an associative nucleus that is part of the major limbic pathway, and primarily projects information related to emotion and memory to the cingulate gyrus.\(^{36}\) Similarly, the DM is involved in emotion\(^{37}\) and cognition, and is connected to the amygdala\(^{38}\) and the anterior cingulate and prefrontal cortices.\(^{36,39,40}\) The VA, which, in combination with the VL, is known as the “motor thalamus,” primarily relays somatomotor information to the supplementary motor area from the basal ganglia. In functional imaging studies of PD, the supplementary motor area is often hypoactive during movement tasks compared with controls.\(^{41–43}\) Taken together, damage to these different thalamic projection fibers may result in widespread impairments\(^{38,43}\) and may contribute to symptom presentation in PD.

Although ex vivo studies remain the criterion standard for pathologic confirmation of PD, it is important to study the brain in vivo to gain insight into the structures affected early in the disease and to evaluate differences across disease stages. As a technique, DTI has proved useful in identifying structural abnormalities in the SN of patients with PD in vivo.\(^{14,15,17,24,44}\) However, DTI studies comparing the thalamus between patients with PD and controls have been less conclusive. In a large cohort, Chan et al\(^{24}\) did not observe differences in FA values at 1.5T between mild to severe patients with PD on antiparkinsonian medication and controls in a ROI that covered a single section of the thalamus. Similarly, there were no differences in mean diffusivity values (FA values were not reported) at 1.5T in a circular ROI placed on section of the thalamus between early-stage medicated patients with PD and controls.\(^{45}\) However, in another large cohort, Péran et al\(^{15}\) reported reduced FA and mean diffusivity values at 3T in the whole thalamus of early stage, but medicated, patients with PD and controls. Most recently, a DTI study that examined the thalamus using both voxelwise and ROI-based approaches reported reduced FA values at 4T in mild to severe medicated patients with PD versus controls.\(^{15}\) Collectively, these studies suggest that magnet field strength and/or ROI size and placement may affect our ability to detect microstructural changes in the thalamus.

As discussed earlier, the nuclei of the thalamus have specialized connectivity and functions, and thus may be differentially affected in disease states, as supported by postmortem research. By studying the thalamus as a single entity, it is unclear whether significant effects in large ROIs or the whole thalamus are driven by degeneration in a few select nuclei, and whether nonsignificant effects are the result of placing small ROIs on relatively spared nuclei. The present study addressed these concerns using high-resolution fiber tracking at 3T, allowing for specific thalamic nuclei to be studied with great sensitiv-
ity. Indeed, we showed that FA values were reduced in the AN, VA, and DM in the patients with PD compared with controls. There was also a marginally significant reduction in FA values from the VL projections of the patients with PD.

Although DTI has proven useful in studying subregions of the SN in PD,\(^4\) relatively few studies have used this technique to examine individual thalamic nuclei. In fact, to date, only the DM and VL have been examined in PD. Consistent with the role of the DM in affect and cognition, Li et al.\(^25\) reported that patients with PD with depression had reduced FA values compared with nondepressed patients with PD, and that these FA values correlated negatively with depression severity. Given that depression is prevalent in PD,\(^46\) it is possible that the patients in our study had greater affective disturbances than the controls. The VL segment of the motor thalamus receives input from the cerebellum\(^47\) and basal ganglia and projects primarily to the pre- and primary motor cortices, retaining its topographic organization throughout. Two previous MR imaging studies at 1.5T have examined the VL in PD, neither of which showed differences in DTI measures between the PD and control groups.\(^44,48\) In the present study, the difference in FA values between the 2 groups approached significance in the VL. Using a high field strength scanner to test larger samples may increase the ability to detect microstructural changes in the VL.

There were 2 thalamic regions with FA values that did not differ significantly between the groups, namely, the VPL/VPM and PU, both of which are sensory nuclei. Human DTI tractography has shown that the PU is connected ipsilaterally to several cortical and subcortical areas, particularly those in the visual pathways.\(^49\) Although its function is not completely understood, the PU is thought to play an important role in higher order visual processing and integration, including visual attention and oculomotor behavior.\(^50\) The VPL and VPM relay touch and proprioceptive information from the contralateral body and head, respectively, to the primary and secondary somatosensory cortices.\(^50\) The fact that we did not show a significant group difference in the PU or VPL/VPM suggests that visual and somatosensory processing should be relatively intact in early-stage PD.

There are a few limitations to the present study. First, a detailed neuropsychologic examination was not performed on the patients with PD. Given that cognitive and affective problems occur frequently in PD,\(^57\) and that we found differences in thalamic nuclei known to be involved in emotion and cognitive processing, it is important to collect these data in the future to better understand the relationship between thalamic degeneration and these nonmotor functions in PD. Second, the reduction of FA values in the VL of patients with PD did not quite reach significance, suggesting that larger sample sizes or more impaired patients may be necessary to detect a difference in this nucleus. Third, intralaminar and midline nuclei (eg, centromedian-parafascicular complex) were not examined because their boundaries could not be distinguished on the MR imaging scans with acceptable interrater reliability. These nuclei, which are involved in arousal and awareness,\(^53\) have shown severe pathology in postmortem studies of PD.\(^1,12,21–23\) Fourth, data in superior cortical areas were not collected due to limitations in the number of sections that can be collected within a reasonable acquisition time. This limited quantification of the tract projections.

**CONCLUSIONS**

The present study provides the first in vivo evidence that there is a disruption of the projection fibers in certain thalamic nuclei, namely the AN, VA, and DM, in de novo patients with PD. These results suggest that structural degeneration occurs beyond the SN in early PD and thus help further our understanding of the neural structures affected by the disease. Specifically, we showed that nuclei involved in motor, cognitive, and affective processes are disrupted, whereas those involved in sensory processes are relatively spared. In addition, this study showed that the current methodology has utility in examining the microstructural integrity of thalamic nuclei in not only traumatic brain injury\(^26\) but also PD. Future research should use these methods longitudinally to determine whether thalamic fiber integrity changes with disease progression and whether it is affected by antiparkinsonian medication. Further, a detailed neuropsychologic examination should be conducted along with DTI of thalamic regions to better understand the impact of fiber degeneration on cognitive, affective, and behavioral functioning in PD.

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