Fixel-Based Analysis of White Matter Degeneration in Patients With Progressive Supranuclear Palsy or Multiple System Atrophy, as Compared to Parkinson’s Disease

Thanh-Thao Nguyen1†, Jur-Shan Cheng2,3,4†, Yao-Liang Chen5,6†, Yu-Chun Lin5,7, Chih-Chien Tsai8, Chih-Song Lu8,10,11, Yi-Hsin Weng10,11,12, Yi-Ming Wu5,6,7, Ngoc-Thanh Hoang1 and Jiun-Jie Wang5,7,8,13

1Department of Radiology, Hue University of Medicine and Pharmacy, Hue University, Hue, Vietnam, 2Clinical Informatics and Medical Statistics Research Center, College of Medicine, Chang Gung University, Taoyuan, Taiwan, 3Department of Biomedical Sciences, College of Medicine, Chang Gung University, Taoyuan, Taiwan, 4Department of Emergency Medicine, Chang Gung Memorial Hospital, Keelung, Taiwan, 5Department of Diagnostic Radiology, Chang Gung Memorial Hospital, Keelung, Taiwan, 6Department of Medical Imaging and Intervention, Chang Gung Memorial Hospital, Linkou, Taiwan, 7Department of Medical Imaging and Radiological Sciences, Chang Gung University, Taoyuan, Taiwan, 8Healthy Aging Research Center, Chang Gung University, Taoyuan, Taiwan, 9Professor Lu Neurological Clinic, Taoyuan, Taiwan, 10Division of Movement Disorders, Department of Neurology, Chang Gung Memorial Hospital, Linkou, Taiwan, 11Neuroscience Research Center, Chang Gung Memorial Hospital, Linkou, Taiwan, 12School of Medicine, Chang Gung University, Taoyuan, Taiwan, 13Medical Imaging Research Center, Institute for Radiological Research, Chang Gung University/Chang Gung Memorial Hospital, Linkou, Taiwan

Introduction: White matter degeneration may contribute to clinical symptoms of parkinsonism.

Objective: We used fixel-based analysis (FBA) to compare the extent and patterns of white matter degeneration in different parkinsonian syndromes—including idiopathic Parkinson’s disease (PD), multiple system atrophy (MSA), and progressive supranuclear palsy (PSP).

Methods: This is a retrospective interpretation of prospectively acquired data of patients recruited in previous studies during 2008 and 2019. Diffusion-weighted images were acquired on a 3-Tesla scanner (diffusion weighting b = 1000 s/mm²—applied along either 64 or 30 non-collinear directions) from 53 patients with PD (men/women: 29/24; mean age: 65.06 ± 5.51 years), 47 with MSA (men/women: 20/27; mean age: 63.00 ± 7.19 years), and 50 with PSP (men/women: 20/30; mean age: 65.96 ± 3.14 years). Non-parametric permutation tests were used to detect intergroup differences in fixel-related indices—including fiber density, fiber cross-section, and their combination.

Results: Patterns of white matter degeneration were significantly different between PD and atypical parkinsonisms (MSA and PSP). Compared with patients with PD, those with MSA and PSP showed a more extensive white matter involvement—noticeably descending tracts from primary motor cortex to corona radiata and cerebral peduncle. Lesions of corpus callosum were specific to PSP and absent in both MSA and PD.
INTRODUCTION

Parkinsonism is a progressive neurodegenerative disorder characterized by resting tremor, rigidity, bradykinesia/akinesia, and postural instability (McFarland, 2016). Although idiopathic Parkinson’s disease (PD) is the most common form of parkinsonism, this condition also comprises multiple system atrophy (MSA) and progressive supranuclear palsy (PSP)—two atypical parkinsonian syndromes that tend to have a more rapid functional deterioration compared with idiopathic PD (McFarland and Hess, 2017). Apart from the pathogenetic changes in basal ganglia, neuroimaging studies in PD have shown various extents of atrophy affecting different brain regions—including a reduced gray matter volume in the frontal (Burton et al., 2004), and pre-frontal lobes (Biundo et al., 2011). Atrophy in the cortical and subcortical areas and noticeably in the cerebellum has been reported in patients with PSP (Giordano et al., 2013), whereas cerebellar white matter atrophy has been described in MSA (Matsusue et al., 2009). Extensive involvement of different brain regions is accompanied by white matter degeneration, which may result in a clinically relevant functional decline (Whitwell et al., 2011).

Although diffusion tensor imaging (DTI) has been previously applied to investigate white matter changes in patients with parkinsonism, this technique suffers from several shortcomings. Errorneous interpretations in DTI may result from oversimplification of the underlying anatomical structures (Mori and Zhang, 2006; Chen et al., 2019), especially in regions with crossing fibers (Jbabdi et al., 2010). In this scenario, fixel-based analysis (FBA) has emerged as a novel approach based on a higher-order diffusion model to compute fiber orientation density function (Raffelt et al., 2017). FBA allows investigating the micro- and macrostructural properties of individual fiber populations within each voxel—with a single fiber population within a voxel termed fixel (Pecheva et al., 2019). Three fixel-based indices can be derived, which are fiber density (FD) —the volume of intra-axonal space of particular fixel; fiber cross-section (FC)—the cross-sectional area of particular fixel; and the combination of fiber density and cross-section (FDC) (Pecheva et al., 2019).

Previous studies supported the clinical usefulness of FBA in investigating white matter degeneration in patients with neurodegenerative diseases (Mito et al., 2018; Rau et al., 2019). Understandings of white matter changes in parkinsonism may provide new insights into the pathogenesis of motor and non-motor symptoms in these clinical entities, which may eventually lead to more personalized care in terms of diagnosis and treatment. The study is original by using FBA, which is a novel development and could provide new interpretation of the underlying changes in the brain as measured by diffusion. We therefore designed this retrospective analysis of prospectively collected data. We used fixel-based analysis to compare the extent and patterns of white matter degeneration in different parkinsonian syndromes—including idiopathic Parkinson’s disease, multiple system atrophy, and progressive supranuclear palsy.

MATERIALS AND METHODS

This is a retrospective interpretation of prospectively acquired data of diffusion MRI. Both diffusion tensor imaging and structural images (as acquired from T1 weighted MPRAGE sequence) were extracted during the period of 2008 and 2019 from medical records. The study protocol complied with the tenets of the Helsinki declaration, and ethical approval was granted by the Chang Gung Medical Foundation Institutional Review Board. Owing to the retrospective nature of the study, the need for informed consent was waived.

Patients

Images were obtained from 53 patients with PD (29 men and 24 women; mean age: 65.06 ± 5.51 years), 47 with MSA (20

TABLE 1 | Demographic and clinical variables.

|               | PD       | MSA      | PSP      | p-value  |
|---------------|----------|----------|----------|----------|
| Number of subject | 53       | 47       | 50       |          |
| Sex (M/F)     | 29/24    | 20/27    | 20/30    | 0.277    |
| Age (years)   | 65.1 ± 5.5 | 63.0 ± 7.2 | 66.0 ± 3.1 | 0.069    |
| Duration (months) | 77.9 ± 52.5a | 49.3 ± 31.4 | 62.9 ± 37.7 | 0.004    |
| MHY           |          |          |          | <0.001   |
| 1             | 15       | 0        | 0        |          |
| 1.5           | 4        | 0        | 1        |          |
| 2             | 8        | 1        | 1        |          |
| 2.5           | 10       | 2        | 2        |          |
| ≥3            | 12       | 29       | 46       |          |
| UPDRS III     | 22.8 ± 15.2abc | 37.2 ± 17.2 | 37.3 ± 17.8 | <0.001   |
| N/A           | 4        | 15       | 0        |          |

Data was not available in both MHY and UPDRS III. PD, Idiopathic Parkinson’s disease; MSA, Multiple system atrophy; PSP, Progressive supranuclear palsy; MHY, Modified Hoehn and Yahr staging scale; UPDRS III, Motor subscate of Unified Parkinson’s Disease Rating Scale.

Discussion: FBA identified specific patterns of white matter changes in MSA and PSP patients compared to PD. Our results proved the utility of FBA in evaluation of implied biological processes of white matter changes in parkinsonism. Our study set the stage for future applications of this technique in patients with parkinsoninian syndromes.

Keywords: fixel-based analysis, white matter, Parkinson’s disease, multiple system atrophy, progressive supranuclear palsy, diffusion weighted imaging
men and 27 women; mean age: 63.00 ± 7.19 years), and 50 with PSP (20 men and 30 women, mean age: 65.96 ± 3.14 years). Eighty-five participants were involved in prior publications, which focused on the analysis of diffusion tensor imaging and its application in the diagnosis and prognosis of patients with Parkinsonism, as well as longitudinal FBA analysis in patients with PD (Wang et al., 2010, 2011; Wai et al., 2012; Lu et al., 2016; Rau et al., 2019b; Chen et al., 2020; Tsai et al., 2020). The current submission employed patients with PD, MSA, and PSP, which investigated white matter difference by FBA analysis. A list of references was added in the Supplementary Table 1. Patients were clinically diagnosed made by two senior neurologists (CSL and YHW, 28 and 21 years of experience, respectively) according to commonly accepted criteria for PD (Hughes et al., 1992), MSA (Gilman et al., 1999), and PSP (Litvan et al., 1996), respectively. The acquisition should be consisted of both diffusion tensor imaging and high-resolution T1-weighted anatomical images. 99mTc-TRODAT-1, which binds to the dopamine transporter, was used to image the dopaminergic system in all participants. Exclusion criteria were as follows: (1) moderate-to-severe dementia; (2) severe dyskinesia; (3) major systemic medical conditions; (4) documented brain abnormalities based on MRI or 18FDG PET findings; (5) history of intracranial surgery; (6) significant neuropsychiatric disorders established by the corresponding diagnostic criteria, including stroke, brain tumor, demyelinating diseases of central nervous system (CNS), major depression, schizophrenia, and Alzheimer’s disease; and (7) pharmacotherapy lasting for >10 years or treatment with drugs capable of crossing the blood-brain-barrier (the only exception being medications for parkinsonian syndromes). The Modified Hoehn and Yahr Staging (Goetz et al., 2004) and the motor subscale of Unified Parkinson Disease Rating Scale (UPDRS III) (Martinez-Martin et al., 1994) were used to assess clinical severity.

FIGURE 1 | Differences in fixel-related indices in PSP compared to PD (column a), MSA compared to PD (column b) and PSP compared to MSA (column c). The involved regions included corona radiata (white arrows in column a and column b) and cerebral peduncle (dashed arrows in columns a, b). Corpus callosum is specifically involved when compared PSP to PD (arrowhead in column a). In PSP group when compared to MSA (column c), the regions with reduced FDC and FD were mainly located in corpus callosum and regions with reduced FC can be found in midbrain and cerebral peduncles (blue). In MSA group when compared to PSP (column c), the regions with reduced FDC, FD and FC were located in middle cerebellar peduncles (red). The streamline showed $p$-values (family-wise error corrected $p < 0.05$, indicated by the colorbar).

FIGURE 2 | Fixel-related indices in patients with PSP compared to that with PD. Reduction of FDC in the main white matter tracts, from superior region of corona radiata (upper row, white arrows), corpus callosum (upper row, black arrow) to posterior limbs of internal capsules (upper row, arrowhead) and cerebral peduncles (upper row, arrowhead). Reduced FD was identified in corona radiata (middle row, white arrows), corpus callosum (middle row, black arrow) and the fornix (middle row, black curved arrows), which were symmetrical in the supratentorial compartment. Reduced FC was noticed in corona radiata to posterior limbs of internal capsules (bottom row, white curved arrow) and cerebral peduncles (bottom row, white arrowhead). The streamline showed $p$-values (family-wise error corrected $p < 0.05$, indicated by the colorbar).
TABLE 2 | Comparison of fiber tracts between patients with PSP and PD.

| Fixel-related indices | Name of tracts | Cluster size | Peak p-value |
|-----------------------|----------------|--------------|--------------|
| FDC                   | Body of corpus callosum | 2532 | 0.00020 |
|                       | Posterior limb of internal capsule L | 1025 | 0.00020 |
|                       | Superior corona radiata L | 992 | 0.00040 |
|                       | Posterior limb of internal capsule R | 896 | 0.00020 |
|                       | Superior corona radiata R | 743 | 0.00040 |
|                       | Cerebral peduncle R | 658 | 0.00020 |
|                       | Cerebral peduncle L | 643 | 0.00020 |
|                       | Anterior limb of internal capsule R | 259 | 0.00040 |
|                       | Superior cerebellar peduncle L | 251 | 0.00020 |
|                       | Superior cerebellar peduncle R | 248 | 0.00020 |
|                       | Corticospinal tract L | 204 | 0.00180 |
|                       | Pontine crossing tract | 190 | 0.00020 |
|                       | Corticospinal tract R | 137 | 0.00180 |
|                       | Medial lemniscus L | 86 | 0.00020 |
|                       | Medial lemniscus R | 63 | 0.00040 |
|                       | Anterior limb of internal capsule L | 54 | 0.01440 |
|                       | External capsule R | 39 | 0.02460 |
|                       | Crura of fornic L | 31 | 0.02620 |
| FD                    | Body of corpus callosum | 3130 | 0.00020 |
|                       | Superior corona radiata R | 489 | 0.00080 |
|                       | Superior corona radiata L | 271 | 0.00100 |
|                       | Crura of fornic R | 171 | 0.00160 |
|                       | Superior cerebellar peduncle R | 162 | 0.00200 |
|                       | Crura of fornic L | 152 | 0.00720 |
|                       | Superior cerebellar peduncle L | 88 | 0.00300 |
|                       | Cerebral peduncle L | 82 | 0.01700 |
|                       | Cerebral peduncle R | 78 | 0.02600 |
|                       | Anterior limb of internal capsule R | 63 | 0.00500 |
|                       | Posterior limb of internal capsule L | 55 | 0.02100 |
|                       | Genu of corpus callosum | 33 | 0.02920 |
|                       | Posterior limb of internal capsule R | 27 | 0.02400 |
|                       | Medial lemniscus R | 21 | 0.01200 |
|                       | Corticospinal tract L | 14 | 0.03300 |
|                       | Corticospinal tract L | 14 | 0.02100 |
|                       | Medial lemniscus L | 11 | 0.02800 |
| FC                    | Superior corona radiata L | 1549 | 0.00020 |
|                       | Posterior limb of internal capsule L | 1194 | 0.00020 |
|                       | Posterior limb of internal capsule R | 1086 | 0.00820 |
|                       | Cerebral peduncle L | 611 | 0.00020 |
|                       | Cerebral peduncle R | 564 | 0.00020 |
|                       | Superior corona radiata R | 552 | 0.00020 |
|                       | Superior cerebellar peduncle L | 261 | 0.00200 |
|                       | Body of corpus callosum | 258 | 0.00040 |
|                       | Pontine crossing tract | 235 | 0.00080 |
|                       | Corticospinal tract L | 234 | 0.00500 |
|                       | superior cerebellar peduncle R | 231 | 0.00020 |
|                       | posterior corona radiata L | 131 | 0.00340 |
|                       | anterior limb of internal capsule R | 127 | 0.00020 |
|                       | corticospinal tract R | 109 | 0.00500 |
|                       | middle cerebellar peduncle | 103 | 0.02200 |

(Continued)

TABLE 2 | Continued

| Fixel-related indices | Name of tracts | Cluster size | Peak p-value |
|-----------------------|----------------|--------------|--------------|
|                       | medial lemniscus L | 82 | 0.00020 |
|                       | medial lemniscus R | 58 | 0.00060 |
|                       | inferior cerebellar peduncle L | 23 | 0.00200 |
|                       | anterior limb of internal capsule L | 11 | 0.00140 |
|                       | inferior cerebellar peduncle R | 10 | 0.00020 |

Two neuro-radiologists (YLC and YMW, 21 and 11 years of experience, respectively), who were blinded to the diagnosis, read the structural MR images independently.

Data Acquisition
All images were acquired on a 3.0-Tesla scanner (Trio Magnetom; Siemens, Erlangen, Germany) using a 12-channel head coil. Diffusion-weighted imaging was performed using two protocols characterized by a different diffusion-sensitive spin-echo EPI sequence. In brief, images with a diffusion weighting b = 1000 s/mm$^2$ were acquired along 64 or 30 non-collinear directions. The voxel size was either $2 \times 2 \times 2$ mm$^3$ or $2 \times 2 \times 3$ mm$^3$. High-resolution T1-weighted anatomical images were obtained using magnetization-prepared rapid gradient-echo (MPRAGE) sequences with the following parameters: TR, 2000 ms; inversion time (TI), 900 ms; TE, 2.63 ms; voxel size, 1 × 1 × 1 mm$^3$.

Image Processing
FBA was carried out on diffusion MRI using MRtrix 3.0 following the recommendations of Raffelt et al. (2017). Pre-processing included denoising by principal component analysis (Veraart et al., 2016), removal of Gibbs ringing (Kellner et al., 2016), as well as correction for motion, distortion, and bias field (Tustison et al., 2010; Andersson and Sotiropoulos, 2015, 2016). Multi-tissue constrained spherical deconvolution was used to estimate fiber orientations distribution in each voxel (Jeurissen et al., 2014). A study-specific template was created by spatial normalization in all of the study patients using symmetric non-linear transformation fiber orientations distribution-based registration. Fiber density and fiber bundle cross-section were calculated within each voxel. A combined measure of fiber density and cross-section (FDC) was computed by multiplying FD by FC (10). For comparison, both Fractional Anisotropy (FA) and Mean Diffusivity (MD) from diffusion tensor imaging were analyzed by using Tract Based Spatial Statistics (TBSS) (Smith et al., 2006) following the recommended procedure.

Statistical Analysis
Differences in fixel-related indices between the three study groups were assessed using non-parametric permutation testing and connectivity-based fixel enhancement as implemented in MRtrix 3.0. Age, sex, and different imaging protocol was used as potential confounding factors. A family-wise error corrected $p$-value $< 0.05$ was considered statistically significant (Nichols and Holmes, 2002).
RESULTS

Demographic and Clinical Variables

Table 1 depicts the general characteristics of the study patients. Age and sex did not differ significantly among the three study groups. Disease duration was longer in patients with PD than in those with MSA ($p = 0.001$). UPDRS III scores were significantly higher in the two atypical parkinsonian syndromes than in patients with PD ($p < 0.001$).

Fixel-Related Indices in the Three Study Groups

Figure 1 displays the changes in fixel-related indices in patients with atypical parkinsonian syndromes (column a: PSP; column b: MSA) compared with PD. All indices were consistently reduced. A 3D visualization of the affected regions with the color encoded the direction of the major fiber bundles are shown in Supplementary Figure 1 (PSP) and 2 (MSA), respectively. With respect to PD, atypical parkinsonian syndromes were characterized by a significant involvement of white matter tract from corona radiata (white arrows in columns a and b) to cerebral peduncle (dashed arrows in columns a and b). Body of corpus callosum was found to be affected when PSP was compared with PD (arrow head in column a).

A comparison of PSP and MSA revealed a differential involvement of various brain areas (column c). Supplementary Figure 3 showed the 3D visualization of the affected regions (color blue: reduction in PSP; color red: reduction in MSA). Regions characterized by reduced FDC (upper row) and FD (middle row) in patients with PSP were chiefly located in the body of corpus callosum, whereas midbrain and cerebral peduncles showed a reduction in FC (blue, bottom row). Regions displaying reduced FDC, FD, and FC values in patients with MSA were localized in the superior cerebellar peduncles (red).

Comparison of Patients With PSP and PD

Figure 2 compares fixel-related indices of patients with PSP and PD. Reductions of FDC (upper row) were evident in the main descending white matter tracts from the superior region of corona radiata (from motor cortex) (white arrows) through body of corpus callosum (black arrow) to posterior limbs of internal capsules (dashed arrows) and into the cerebral peduncles (arrow head). The regions with changes are consistent with that of reduced FD (middle row), noticeably in corona radiata (white arrows), and corpus callosum (black arrow). There was involvement of the medial thalamus (black curved arrow)—with a symmetrical involvement of the supratentorial compartment. Similar regions of reductions in FC (bottom row) were evident from corona radiata to the posterior limbs of internal capsules and cerebral peduncles (arrow head). Regions with a concomitant reduction of both FD and FC were generally overlapping to those with a reduced FDC. However, an isolated decreased of FD was evident in the thalamus. The names of tracts with significant difference, together with the cluster sizes and the peak $p$-values were summarized in Table 2.

Comparison of Patients With MSA and PD

Figure 3 compares fixel-related indices of patients with MSA and PD. Reductions of FDC (top row) were evident in the main descending white matter pathways from the left corona radiata (white arrows) to bilateral posterior limbs of internal capsule (white dashed arrows), cerebral peduncles, transverse pontine fibers (white arrow head), and bilateral middle cerebellar peduncles (black arrows). FD was found to be significantly reduced (middle row) in bilateral cerebral peduncles (white arrows) and middle cerebellar peduncles (black arrows).
Interestingly, regions with reduced FC (bottom row) almost invariably overlap with those showing a reduction in FDC. The names of tracts with significant difference, together with the cluster sizes and the peak \( p \)-values were summarized in Table 3.

### Comparison of Patients With PSP and MSA

Figure 4 compares fixel-related indices of patients with PSP and MSA. Reductions of FDC and FD in patients with PSP were evident in body of corpus callosum (white arrows), whereas FC was found to be lowered in the midbrain (white dashed arrows). All fixel-related indices (FDC, FD, and FC) of patients with MSA were reduced in middle cerebellar peduncles (Figure 4, black arrows). The names of tracts with significant difference, together with the cluster sizes and the peak \( p \)-values were summarized in Table 4.

### Tract-Based Spatial Statistics in Patients With PSP or MSA, as Compared to PD

For comparison to conventional diffusion tensor imaging, Figure 5 shows the result from tract-based spatial statistics relative to patients with PD. In patients with PSP, reduced FA (panel a) and increased MD (panel b) can be found in bilateral corona radiata, internal capsules, superior longitudinal fasciculi, posterior thalamic radiation, as well as genu and splenium of the corpus callosum. In patients with MSA, only increased MD (panel c) was found, which was located in the bilateral cerebellar peduncle.

### DISCUSSION

#### Main Findings

FBA was used to assess the patterns of white matter changes in three different forms of parkinsonism. Our findings revealed profound differences in terms of white matter involvement among groups—with affected regions being more extensive in MSA and PSP than in PD. These results are consistent with the notion that both PSP and MSA portend a higher burden of clinical disability. Further, brain regions affected by white matter changes were in accordance with the current models of disease pathogenesis (Boxer et al., 2017; Meissner et al., 2019). Our analysis method might provide an interesting insight into the pathophysiology of various parkinsonisms. Taken together, our data indicate that the distribution and severity of white matter involvement in patients with PSP and MSA are characteristically different.

The identified regions can be overlapped between FBA and conventional DTI analysis using TBSS. Both techniques might provide complementary information regarding to the underlying microstructural changes. However, because different post-processing procedures may lead to either false-positive or false-negative results (Kuchling et al., 2018), the choice of the analysis approach might depend on the research question of interest. Our study demonstrated the use of fixel-related indices as potential biomarkers of disease progression. It may set the stage for future applications of this technique in patients with parkinsonian syndromes.

Table 3: Comparison of fiber tracts between patients with MSA and PD.

| Fixel-related indices | Name of tracts                  | Cluster size | Peak \( p \)-value |
|-----------------------|--------------------------------|--------------|-------------------|
| FDC                   | Posterior limb of internal capsule L | 1027         | 0.00020           |
|                       | Posterior limb of internal capsule R | 931          | 0.00020           |
|                       | Middle cerebellar peduncle       | 800          | 0.00020           |
|                       | Cerebral peduncle R              | 768          | 0.00020           |
|                       | Cerebral peduncle L              | 701          | 0.00020           |
|                       | Superior corona radiata L        | 378          | 0.00760           |
|                       | Corticospinal tract L            | 321          | 0.00020           |
|                       | Corticospinal tract R            | 305          | 0.00020           |
|                       | Pontine crossing tract           | 293          | 0.00020           |
|                       | Superior cerebellar peduncle R   | 178          | 0.00018           |
|                       | Superior cerebellar peduncle L   | 149          | 0.00060           |
|                       | Anterior limb of internal capsule R | 125          | 0.00060           |
|                       | Medial lemniscus L               | 83           | 0.00320           |
|                       | Medial lemniscus R               | 58           | 0.00020           |
|                       | Retrolenticular part of internal capsule L | 52       | 0.00080           |
|                       | Posterior corona radiata L       | 52           | 0.02280           |
|                       | Inferior cerebellar peduncle L   | 49           | 0.00120           |
|                       | Retrolenticular part of internal capsule R | 49       | 0.00180           |
|                       | Anterior limb of internal capsule L | 29           | 0.01390           |
|                       | Superior corona radiata R        | 14           | 0.01500           |
|                       | Middle cerebellar peduncle       | 411          | 0.00020           |
|                       | Cerebral peduncle R              | 368          | 0.00020           |
|                       | Cerebral peduncle L              | 273          | 0.00040           |
|                       | Posterior limb of internal capsule R | 192          | 0.00060           |
|                       | Corticospinal tract L            | 171          | 0.00040           |
|                       | Posterior limb of internal capsule L | 135          | 0.000620          |
|                       | Corticospinal tract R            | 132          | 0.00140           |
|                       | Pontine crossing tract           | 67           | 0.00640           |
|                       | Retrolenticular part of internal capsule R | 29       | 0.00060           |
|                       | Superior part_cingulum_R         | 27           | 0.02400           |
|                       | Superior cerebellar peduncle R   | 24           | 0.02000           |
|                       | Crura of fornix L                | 17           | 0.03800           |
|                       | Crura of fornix R                | 14           | 0.02900           |
| FD                    | Posterior limb of internal capsule L | 1139         | 0.00020           |
|                       | Posterior limb of internal capsule R | 1024         | 0.00020           |
|                       | Superior corona radiata L        | 991          | 0.00100           |
|                       | Middle cerebellar peduncle       | 792          | 0.00020           |
|                       | Cerebral peduncle R              | 662          | 0.00020           |
|                       | Cerebral peduncle L              | 633          | 0.00020           |
|                       | Corticospinal tract L            | 334          | 0.00020           |
|                       | Corticospinal tract R            | 326          | 0.00020           |
|                       | Pontine crossing tract           | 269          | 0.00020           |
|                       | Superior cerebellar peduncle L   | 124          | 0.00040           |
|                       | Posterior corona radiata L       | 117          | 0.00700           |
|                       | Superior corona radiata R        | 96           | 0.01180           |
|                       | Anterior limb of internal capsule R | 71           | 0.00340           |
|                       | Superior cerebellar peduncle R   | 66           | 0.00480           |
|                       | Inferior cerebellar peduncle L   | 56           | 0.00440           |
|                       | Medial lemniscus L               | 21           | 0.00700           |
|                       | Anterior limb of internal capsule L | 18           | 0.00700           |
FIGURE 4 | Fixel-related indices in patients with PSP and MSA. Reduced FDC and FD were noticeable in corpus callosum (white arrows), while reduced FC were found in midbrain (white dashed arrows) in patients with PSP. On the other hand, the regions with reduced FDC, FD and FC were found in middle cerebellar peduncles (black arrows) in patients with MSA. The streamline showed p-values (family-wise error corrected $p < 0.05$, indicated by the colorbar).

TABLE 4 | Comparison of fiber tracts between patients with PSP and MSA.

| Fixel-related indices | Name of tract             | Cluster size | Peak p-value |
|-----------------------|---------------------------|--------------|--------------|
| FDC                   | Body of corpus callosum   | 1048         | 0.01360      |
|                       | Middle cerebellar peduncle| 118          | 0.03000      |
|                       | Superior corona radiata R | 110          | 0.01340      |
|                       | Corticospinal tract L     | 40           | 0.02140      |
|                       | Superior corona radiata L | 35           | 0.01700      |
|                       | Corticospinal tract R     | 25           | 0.02740      |
|                       | Cerebral peduncle L       | 11           | 0.02400      |
| FD                    | Body of corpus callosum   | 1196         | 0.00640      |
|                       | Superior corona radiata R | 55           | 0.01040      |
|                       | Superior corona radiata L | 22           | 0.02940      |
|                       | Corticospinal tract L     | 18           | 0.02680      |
|                       | Cerebral peduncle L       | 17           | 0.02000      |
|                       | Middle cerebellar peduncle| 13           | 0.03260      |
|                       | Corticospinal tract R     | 10           | 0.03080      |
| FC                    | Middle cerebellar peduncle| 58           | 0.03160      |
|                       | Superior cerebellar peduncle R | 48       | 0.01680      |
|                       | Superior cerebellar peduncle L | 45     | 0.01340      |
|                       | Medial lemniscus L        | 38           | 0.01080      |
|                       | Medial lemniscus R        | 32           | 0.01680      |

PSP: From Corpus Callosum, Internal Capsules, Thalamic Radiation to Midbrain

The corpus callosum consists of white matter fibers that interconnect the motor areas of the two hemispheres. When compared to patients with PD, reductions of FDC in the corpus callosum of PSP overlapped considerably with FD declines. Similarly, when compared to patients with MSA, a reduced FDC in the corpus callosum can be noticed, a change which was chiefly driven by a decline in FD rather than in FC. Reductions of FD may be related to a lowered intra-axonal volume as a result of axonal degeneration (Raffelt et al., 2017).

In this scenario, our findings might reflect the occurrence of Wallerian degeneration in the affected descending pathways—possibly secondary to cortical gray matter atrophy (Worker et al., 2014). Our findings support the early-stage white matter involvement in corpus callosum in PSP, which may aid in our understanding of the difference in pathogenesis between PSP and MSA.

Previous studies have reported that patients with PSP display abnormal diffusion metrics in the main white matter tract in addition to corpus callosum—for example, the internal capsule (Padovani et al., 2006; Agosta et al., 2012). Our findings of altered fixel-related indices in these regions suggest that the previously reported abnormalities are likely the results of microstructural white matter injury. Patients with PSP have a reduced FDC in both the corona radiata and internal capsules. Such alterations are in line with the higher severity of motor deficits observed in patients with PSP than in those with PD (Alster et al., 2020).

Notably, regions with reduced FDC in the corona radiata, internal capsules, and midbrain had broad overlaps with those showing diminished FC. The latter alteration can be attributed to a narrowing of extra-axonal space followed by axonal loss and disruption of myelin sheaths as a result of white matter atrophy (Raffelt et al., 2017). Interestingly, regions with reduced FD was found in bilateral medial thalami. These fibers are parts of thalamic radiation which connect the thalami with the cortex via internal capsule (Sun et al., 2018). Reduced FD, rather than FC, might imply early changes within these fibers. Similar regions have been reported in a quantitative study of histology, which showed vacuolation and glial inclusion in descending white matter tract from frontal cortex through the internal capsules and midbrain (Armstrong, 2013). Our findings suggest the involvement of the thalamic radiation in PSP, which may help distinguishing this disease from PD and MSA.

Taken together, these results suggest that the severity of white matter damage in internal capsules, thalamic radiation and
midbrain in PSP is more pronounced than in PD. Our findings suggest that the previously reported abnormalities are likely the results of microstructural white matter injury.

**MSA: Corona Radiata, Internal Capsule, and Middle Cerebellar Peduncles**

MSA is specifically characterized by an involvement of the middle cerebellar peduncles (Poewe and Wenning, 2002). When patients with MSA were compared to those with PD, we found significant reductions of both FDC and FC in the main descending white matter pathways. A previous DTI study identified large areas characterized by reduced FA and increased MD in white matter tracts—especially in the supratentorial and infratentorial compartments (Tha et al., 2010). Here, we provide evidence of a substantial axonal loss in the white matter located in these areas.

When compared with those with PD as well as PSP, we found significant reductions of FDC in the middle cerebellar peduncles—which was accompanied by parallel declines in both FD and FC. Increased diffusivity has been previously reported in this region (Tsukamoto et al., 2012). Our results indicate that these regions, characterized by a reduced FDC, were driven by a concomitant decline in both FD and FC suggesting that axonal degeneration is likely the main contributor to microstructural damage. MSA can be associated with a more severe axonal neurodegeneration and white matter atrophy in these areas compared with either PD or PSP. Previous postmortem pathology study showed that the white matter atrophy of the cerebellum in patients with MSA can be attributed to the loss of myelinated fibers and gliosis (Matsusue et al., 2009b). Our observation of reduced FD and FC, as explained by the simultaneous presence of axonal loss and white matter atrophy, might ultimately reflect more severe derangements as related to the motor deficits in patients with MSA than in those with PD.

Significant reductions of both FDC and FC were also found in the main descending white matter pathways, namely the left corona radiata, bilateral posterior limbs of internal capsule, cerebral peduncles, and ventral and dorsal transverse pontine fibers. Concomitant reduction of all three fixel-related indices in these regions implies severe white matter damages. This is in line with the more severe morbidity of MSA in comparison to PSP and PD. Here, we provide image-based evidence of a substantial axonal loss in the white matter located in these areas.

**Limitation**

Our findings should be interpreted in the context of some limitations. In this study, patients with PD was used as a “parkinsonian control” group with which both the patients with MSA or PSP were compared to. In the future study when using a prospective design, we will recruit the healthy control in our study for a comprehensive understanding of the white matter involvement in Parkinsonism. Secondly, it may be argued that the study groups were different in terms of imaging protocols. However, the effect from this variable, together with age and sex, has been considered and controlled in the statistical analysis. Provided the acquisition and the analysis pipeline are the same, these measures can be quantifiable to allow for assessment of disease severity within study. However,
it would be necessary to perform advanced statistical analysis to determine the sensitivity and specificity before its potential application in early-stage therapeutic trials, or to monitor disease progression. More radiologic—pathologic correlation studies are necessary to scrutinize the biologic mechanisms of these fixed-related indices’ alteration. These caveats notwithstanding, we successfully applied FBA and identified specific patterns of white matter degeneration that distinguished PD from atypical parkinsonisms (MSA and PSP).

**DATA AVAILABILITY STATEMENT**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by the study protocol complied with the tenets of the Helsinki declaration, and ethical approval was granted by the Chang Gung Medical Foundation Institutional Review Board. Owing to the retrospective nature of the study, the need for informed consent was waived. Written informed consent for participation was not required for this study in accordance with the national legislation and the institutional requirements.

**AUTHOR CONTRIBUTIONS**

T-TN: conception and design of study, statistical analysis, and writing the first draft. Y-LG, Y-CL, C-SL, Y-HW, and Y-MW: acquisition of clinical data and revision the manuscript. C-CT: conception and design of study, writing and revision the manuscript, study supervision, and obtaining funding. All authors contributed to the article and approved the submitted version.

**FUNDING**

This work was supported by the Ministry of Science and Technology Taiwan (grants MOST 106-2314-B-182-018-MY3, MOST 109-2221-E-182-009-MY3, MOST 109-2314-B-182-021-MY3); the Healthy Aging Research Center (grants EMRPD110501, EMRPD110471, EMRPD1K0451, EMRPD1K0481); and the Chang Gung Memorial Hospital (grants CMRPD1G0561-2, CMRPG2B0251, and CMRPG2J0141).

**ACKNOWLEDGMENTS**

The imaging facility was supported by the Imaging Core Laboratory of the Institute for Radiological Research and the Center for Advanced Molecular Imaging and Translation. The authors thank the Neuroscience Research Center (Chang Gung Memorial Hospital) and the Healthy Aging Research Center (Chang Gung University) for their invaluable support.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fnagi.2021.625874/full#supplementary-material

**REFERENCES**

Agosta, F., Pievani, M., Svetel, M., Jecmenica Lukic, M., Copetti, M., Tomic, A., et al. (2012). Diffusion tensor MRI contributes to differentiate Richardson’s syndrome from PSP-parkinsonism. Neurobiol. Aging 33, 2817–2826. doi: 10.1016/j.neurobiolaging.2012.02.002

Altier, P., Madetko, N., Koziorowski, D., and Friedman, A. (2020). Progressive Supranuclear Palsy-Parkinsonism Predominant (PSP-P): a clinical challenge at the boundaries of PSP and Parkinson’s Disease (PD). Front. Neurol. 11:180. doi: 10.3389/fneur.2020.00180

Andersson, J. L., and Sotiropoulos, S. N. (2015). Non-parametric representation and prediction of single- and multi-shell diffusion-weighted MRI data using Gaussian processes. NeuroImage 122, 166–176. doi: 10.1016/j.neuroimage.2015.07.067

Andersson, J. L. R., and Sotiropoulos, S. N. (2016). An integrated approach to correction for off-resonance effects and subject movement in diffusion MR imaging. Neuroimage 125, 1063–1078. doi: 10.1016/j.neuroimage.2015.10.019

Armstrong, R. A. (2013). White matter pathology in progressive supranuclear palsy (PSP): a quantitative study of 8 cases. Clin. Neuropathol. 32, 399–405. doi: 10.5414/NNP00608

Biundo, R., Formento-Dojot, P., Facchini, S., Vallelunga, A., Ghezzo, L., Foscolo, L., et al. (2011). Brain volume changes in Parkinson’s disease and their relationship with cognitive and behavioural abnormalities. J. Neurol. Sci. 310, 64–69. doi: 10.1016/j.jns.2011.08.001

Boxer, A. L., Yu, J. T., Golbe, L. I., Litvan, I., Lang, A. E., and Hoglinger, G. U. (2017). Advances in progressive supranuclear palsy: new diagnostic criteria, biomarkers, and therapeutic approaches. Lancet Neurol. 16, 552–563. doi: 10.1016/S1474-4422(17)30157-6

Burton, E. J., McKeith, I. G., Burn, D. J., Williams, E. D., and O’Brien, J. T. (2004). Cerebral atrophy in Parkinson’s disease with and without dementia: a comparison with Alzheimer’s disease, dementia with Lewy bodies and controls. Brain 127, 791–800. doi: 10.1093/brain/awh088

Chen, Y.-L., Zhao, X.-A., Ng, S.-H., Lu, C.-S., Lin, Y.-C., Cheng, J.-S., et al. (2020). Prediction of the clinical severity of progressive supranuclear palsy by diffusion tensor imaging. J. Clin. Med. 9:40. doi: 10.3390/jcm9010040

Chen, Y.-L., Lin, Y.-J., Lin, S.-H., Tsai, C.-C., Lin, Y.-C., Cheng, J.-S., et al. (2019). The effect of spatial resolution on the reproducibility of diffusion imaging when controlled signal to noise ratio. Biomed. J. 42, 268–276. doi: 10.1016/j.bj.2019.03.002

Gilman, S., Low, P. A., Quinn, N., Albanese, A., Ben-Shlomo, Y., Fowler, C. J., et al. (1999). Consensus statement on the diagnosis of multiple system atrophy. J. Neurol. Sci. 163, 94–98. doi: 10.1016/S0022-5188(99)00304-9

Giordano, A., Tessitore, A., Corbo, D., Cirillo, G., de Micco, R., Russo, A., et al. (2013). Clinical and cognitive correlations of regional gray matter atrophy in progressive supranuclear palsy. Parkinsonism Relat. Disord. 19, 590–594. doi: 10.1016/j.parkreldis.2013.02.005

Goetz, C. G., Poewe, W., Sasaki, A., Jhangiani, S. A., Storlazzi, C. T., Schrag, A. J., et al. (2004). Movement Disorder society task force report on the Hoehn and Yahr staging scale: status and recommendations. Mov. Disord. 19, 1020–1028. doi: 10.1002/mds.20213

Hughes, A. J., Ben-Shlomo, Y., Daniel, S. E., and Lees, A. J. (1992). What features improve the accuracy of clinical diagnosis in Parkinson’s disease:
a clinicopathologic study. *Neurology* 42, 1142–1146. doi: 10.1212/WNL.42.6.1142

Jbabdi, S., Behrens, T. E., and Smith, S. M. (2010). Crossing fibres in tract-based spatial statistics. *Neuroimage* 49, 249–256. doi: 10.1016/j.neuroimage.2009.08.039

Jeuring, B., Tournier, J. D., Dhollander, T., Connelly, A., and Sijbers, J. (2014). Multi-tissue constrained spherical deconvolution for improved analysis of multi-shell diffusion MRI data. *Neuroimage* 103, 411–426. doi: 10.1016/j.neuroimage.2014.07.061

Kellner, E., Dhital, B., Kiselev, V. G., and Reisert, M. (2016). Gibbs-ringging artifact removal based on local subvoxel-shifts. *Magn. Reson. Med.* 76, 1574–1581. doi: 10.1002/mrm.26054

Kuchling, J., Backner, Y., Oertel, F. C., Raz, N., Bellmann-Strobl, J., Connelly, A., and Sijbers, J. (2014). Comparison of probabilistic tractography and tract-based spatial statistics for assessing optic radiation damage in patients with autoimmune inflammatory disorders of the central nervous system. *Neuroimage Clin.* 19, 538–550. doi: 10.1016/j.nicl.2018.05.004

Litvan, I., Agid, Y., Calne, D., Campbell, G., Duvoisin, R. C., et al. (1996). Clinical research criteria for the diagnosis of progressive supranuclear palsy (Steele-Richardson-Olszewski syndrome): report of the NINDS-SPSP international workshop. *Neurology* 47, 1–9. doi: 10.1212/WNL.42.6.1142

McFarland, N. R., and Hess, C. W. (2017). Recognizing atypical Parkinsonisms: future perspectives. *Semin. Neurol.* 37, 1629–1642. doi: 10.1055/s-0037-1602422

Mori, S., and Zhang, J. (2006). Principles of diffusion tensor imaging and its applications to basic neuroscience research. *Neuron* 51, 527–539. doi: 10.1016/neuron.2006.08.012

Nichols, T. E., and Holmes, A. P. (2002). Nonparametric permutation tests for functional neuroimaging: a primer with examples. *Hum. Brain Mapp.* 15, 1–25. doi: 10.1002/hbm.1058

Padovani, A., Borroni, B., Brambati, S. M., Agosti, C., Broli, M., Alonso, R., et al. (2006). Diffusion tensor imaging and voxel based morphometry study in early progressive supranuclear palsy. *J. Neurol. Neurosurg. Psychiatry* 77, 457–463. doi: 10.1136/jnnp.2005.057713

Pechova, D., Tournier, J. D., Pietsch, M., Christiaens, D., Bataille, D., Alexander, D. C., et al. (2019). Fixel-based analysis of the preterm brain: disentangling bundle-specific white matter microstructural and macrostructural changes in relation to clinical risk factors. *Neuroimage Clin.* 23:101820. doi: 10.1016/j.nicl.2019.101820

Poewe, W., and Wenning, G. (2002). The differential diagnosis of Parkinson’s disease. *Eur. J. Neurol.* 9, 23–30. doi: 10.1046/j.1468-1331.9.s3.3.x

Raffelt, D., Tournier, J. D., Smith, R. E., Vaughan, D. N., Jackson, G., Ridgway, G. R., et al. (2017). Investigating white matter fibre density and morphology using fixel-based analysis. *Neuroimage* 144, 58–73. doi: 10.1016/j.neuroimage.2016.09.029

Rau, Y. A., Wang, S. M., Tournier, J. D., Lin, S. H., Lu, C. S., Weng, Y. H., et al. (2019). A longitudinal fixel-based analysis of white matter alterations in patients with Parkinson’s disease. *Neuroimage Clin.* 24:102098. doi: 10.1016/j.nicl.2019.102098

Smith, S. M., Jenkinson, M., Johansen-Berg, H., Rueckert, D., Nichols, T. E., Mackay, C. E., et al. (2006). Tract-based spatial statistics: voxelwise analysis of multi-subject diffusion data. *Neuroimage* 31, 1487–1505. doi: 10.1016/j.neuroimage.2006.02.024

Sun, C., Wang, Y., Cui, R., Wu, C., Li, X., Bao, Y., et al. (2018). Human thalamic-prefrontal peduncle connectivity revealed by diffusion spectrum imaging fiber tracking. *Front. Neuroanat.* 12:24. doi: 10.3389/fnana.2018.00024

Tha, K. K., Terae, S., Yabe, I., Miyamoto, T., Soma, H., Zaitsu, Y., et al. (2010). Microstructural white matter abnormalities of multiple system atrophy: in vivo topographic illustration by using diffusion-tensor MR imaging. *Radiology* 255, 563–569. doi: 10.1148/radiol.10090988

Tsai, C.-C., Weng, Y.-H., Lin, S.-H., Chen, P.-Y., and Wu, Y.-M. (2020). A method for the prediction of clinical outcome using diffusion magnetic resonance imaging: application on Parkinson’s disease. *J. Clin. Med.* 9:647. doi: 10.3390/jcm9030647

Tsukamoto, K., Matsusue, E., Kanasaki, Y., Kakite, S., Fujii, S., Kaminou, T., et al. (2012). Significance of apparent diffusion coefficient measurement for the differential diagnosis of multiple system atrophy, progressive supranuclear palsy, and Parkinson’s disease: evaluation by 3.0-T MR imaging. *Neuroradiology* 54, 947–955. doi: 10.1007/s00234-012-1009-9

Tustison, N. J., Avants, B. B., Cook, P. A., Zheng, Y., Egan, A., Yushkevich, P. A., et al. (2010). N4ITK: improved N3 bias correction. *IEEE Trans. Med. Imaging* 29, 1310–1320. doi: 10.1109/TMI.2010.2046908

Veraart, J., Novikov, D. S., Christiaens, D., Ades-Aron, B., Sijbers, J., and Fieremans, E. (2016). Denoising of diffusion MRI using random matrix theory. *Neuroimage* 142, 394–406. doi: 10.1016/j.neuroimage.2016.08.016

Wai, Y.-Y., Wang, J.-J., Lin, W.-Y., Ma, H.-K., Ng, S.-H., et al. (2012). Cortical involvement in a gait-related imagery task: comparison between Parkinson’s disease and normal aging. *Parkinsonism Relat. Disord.* 18, 537–542. doi: 10.1016/j.parkreldis.2012.02.004

Wang, J., Wai, Y., Lin, W. Y., Ng, S., Wang, C. H., Hsieh, R., et al. (2010). Microstructural changes in patients with progressive supranuclear palsy: a diffusion tensor imaging study. *J. Magn. Reson. Imaging* 32, 69–75. doi: 10.1002/jmri.22229

Wang, J.-J., Lin, W.-Y., Yu, C.-S., Weng, Y.-H., Ng, S.-H., and Cheng, J.-S., Lin, W.-Y., Ma, H.-K., Ng, S.-H., et al. (2011). Parkinson disease: diagnostic utility of diffusion kurtosis imaging. *Radiology* 261, 210–217. doi: 10.1148/radiol.11102277

Whitwell, J. L., Master, A. V., Avula, R., Kantarci, K., Eggers, S. D., Edmonson, H. A., et al. (2011). Clinical correlates of white matter tract degeneration in progressive supranuclear palsy. *Arch. Neurol.* 68, 753–760. doi: 10.1001/archneurol.2011.107

Worker, A., Blain, C., Jarosz, J., Chaudhuri, K. R., Barker, G. J., Williams, S. C., et al. (2014). Diffusion tensor imaging of Parkinson’s disease, multiple system atrophy and progressive supranuclear palsy: a tract-based spatial statistics study. *PLoS ONE* 9:e116238. doi: 10.1371/journal.pone.012638

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2021 Nguyen, Cheng, Chen, Lin, Tsai, Lu, Weng, Wu, Hoang and Wang. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.