Brain Diffusion Imaging and Tractography to Distinguish Clinical Severity of Human PLP1-Related Disorders
Catherine Sarret, Jean-Jacques Lemaire, Anna Sontheimer, Jerome Coste, Nadia Savy, Bruno Pereira, Bastien Roche, Odile Boespflug-Tanguy

To cite this version:
Catherine Sarret, Jean-Jacques Lemaire, Anna Sontheimer, Jerome Coste, Nadia Savy, et al.. Brain Diffusion Imaging and Tractography to Distinguish Clinical Severity of Human PLP1-Related Disorders. Developmental Neuroscience, Karger, 2018, 40 (4), pp.301-311. 10.1159/000492218. hal-01952380

HAL Id: hal-01952380
https://hal.archives-ouvertes.fr/hal-01952380
Submitted on 13 Dec 2018

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Brain diffusion imaging and tractography to distinguish clinical severity of human \textit{PLP1}-related disorders

Authors

Catherine Sarret\textsuperscript{1,2*}, Jean-Jacques Lemaire\textsuperscript{1,3}, Anna Sontheimer\textsuperscript{1,3}, Jérôme Coste\textsuperscript{1,3}, Nadia Savy\textsuperscript{1,2}, Bruno Pereira\textsuperscript{4}, Bastien Roche\textsuperscript{5}, Odile Boespflug-Tanguy\textsuperscript{5,6}.

\textsuperscript{1}Institut Pascal, CNRS, Université Clermont Auvergne, SIGMA, France
\textsuperscript{2}Service de génétique médicale, Centre de compétence des leucodystrophies et leucoencéphalopathies de cause rare, CHU Clermont-Ferrand, France
\textsuperscript{3}Service de Neurochirurgie, CHU Clermont-Ferrand, France
\textsuperscript{4}Unité de Biostatistiques (Délegation Recherche Clinique & Innovation), CHU Clermont-Ferrand, France
\textsuperscript{5}Service de Neurologie Pédiatrique et Maladies Métaboliques, Centre de référence des leucodystrophies et leucoencéphalopathies de cause rare, CHU-APHP Robert-Debré, Paris, France.
\textsuperscript{6}INSERM UMR1141 PROTECT Université Paris-Diderot-Sorbonne Paris Cité, France

*Correspondence: Catherine Sarret, Equipe IGCNC (Institut Pascal, UMR CNRS-UCA-SIGMA), CHU de Clermont-Ferrand, France, 58 rue Montalember, 63003 Clermont-Ferrand Cedex, France. Tel: +33(0)473750296. Fax: +33(0)473751201. E-mail: csarret@chu-clermontferrand.fr.
Abstract

Aims: We performed quantitative diffusion tensor imaging and brain tractography to distinguish clinical severity in a series of 35 patients with hypomyelinating PLP1-related disorders classified using the Motor Developmental Score according to best motor function acquired before 5 years and the gross motor function measure (GMFM) at the time of MRI acquisition.

Methods: We calculated fractional anisotropy and diffusivity values in 26 regions of interest, and the numbers of fibers and volumes of hemisphere tractograms. Fiber bundles on tractograms were characterized according to three criteria: size, direction of main-stream fibers, and connectivity of bundles (extratelencephalic projections, commissural fibers, and intrahemispheric connections).

Results: Age-adjusted multivariate analysis in three severity groups revealed increased isotropic diffusion in the superior cerebellar peduncle and grey matter in the most severe group, and larger tractogram volumes and increased numbers of fibers in the least severe group. Tractogram patterns showed preserved extratelencephalic projections and a main anterior-posterior aspect of intrahemispheric fibers in most patients, whereas interhemispheric connectivity was variable. The most severe and intermediate patients had less intrahemispheric connectivity, with a frequent predominant anterior-posterior direction of main-stream fibers.

Interpretation: Diffusion tensor imaging and tractographic parameters can operate as biomarkers to distinguish clinical severity in PLP1-related disorders and could improve our understanding of hypomyelinating leukodystrophies.

Key-words

PLP1-related disorders, Pelizaeus–Merzbacher disease, Spastic paraplegia type 2, diffusion tensor imaging, tractography, Proteolipid Protein

Running Title: DTI in PLP1-related disorders
Introduction

Pelizaeus-Merzbacher disease (PMD, OMIM312080) and spastic paraplegia type 2 (SPG2, OMIM312920) are X-linked hypomyelinating leukodystrophies caused by mutations in the Proteolipid protein 1 (PLP1) gene that encodes the major central nervous system (CNS) myelin proteins, PLP and DM20 in oligodendrocytes [1,2]. The clinical findings associated with PLP1 mutations span a broad-spectrum continuum extending from severe congenital PMD forms to relatively mild late-onset SPG2, leading to the concept of PLP1-related disorders [3,4]. We previously classified patients according to the best motor function achieved between the ages of 2 and 5 years to demonstrate a genotype–phenotype correlation [3]. PMD1 (head control) and PMD2 (sitting position) are the most frequent forms and are usually associated with PLP1 duplications. Missense mutations are rare but usually cause the most severe forms without motor acquisitions and dystonia (PMD0, also named “congenital” forms). Myelin deficiency is the main histological feature in human and mouse models of these severe forms [5,6]. PLP1 loss-of-function (null mutations and large deletions) leads to the mildest PMD3/SPG2 forms with acquisition of walking capacities with/without aid and relatively preserved cognitive development. Murine models and humans lacking PLP1 are characterized by axonal degeneration with a small enlargement of the interlamellar space of the myelin sheath [2,7]. In all forms, the myelin deficiency is demonstrated by the extensive abnormalities of multimodal evoked potentials and the diffuse hypomyelinated pattern of the supratentorial white matter (WM) magnetic resonance imaging (MRI) [8]. Most patients clinically improve during childhood, exactly like patients with cerebral palsy. Some progression of myelination is observed until 12 years of age [9]. The second decade is marked by neurological deterioration with cortico-subcortical atrophy on MRI, leading to severe quadriplegia, amyotrophy, optic atrophy and cognitive decline in young adults, even in the mildest forms [10]. Several recent studies have underlined the correlation between clinical severity and brain atrophy on MRI [9,11,12].

Using diffusion tensor imaging (DTI), Laukka et al. found that the WM mean and radial diffusivities of 12 PLP1-mutated patients were higher than controls, in accordance with the severe myelin deficiency observed in PMD [13]. Axial diffusivity, which is assumed to be more specific to axon properties, was much less affected, except in the most severe patients. Here we investigated the ability of DTI parameters to distinguish the different severity groups (PMD0-1, PMD2, and PMD3-4) and to correlate with clinical severity in a cohort of 35 PLP1-mutated patients. We extended the WM analysis with tractography in an attempt to identify patterns of WM fiber damage.
Patients and methods

Patients

Thirty-five patients from 32 families with a genetically-confirmed diagnosis of PLP1 mutation (19 duplications, 9 missense mutations and 7 null mutations) were included in this prospective study from January 2004 to December 2009, after first obtaining approval from the institutional review board (CPP Sud-Est VI No. AU788, CNIL No. 1406552, AFSSAPS No. B90298-60) and signed informed consent from the parents. For MRI acquisition, sedation by intrarectal pentobarbital: 3 to 5 mg/kg (maximum dose: 100 mg) and oral alimemazine (4% drops, 1 drop/kg, i.e. 1.25 mg/kg), was performed in patients aged under 6 years or requiring sedation to avoid movement artifacts. Our study focused on a comparison of patients with distinct degrees of disease severity. Patients were classified according to the best motor developmental score (MDS) observed between age 2 and 5 years: PMD0, without motor acquisition (n=4 patients); PMD1, head control (n=5); PMD2, sitting position (n=16); PMD3, walking with support (n=8); PMD4, independent walking (n=2). An 88-item gross motor function measure (GMFM) was assessed for all patients at the time of MRI acquisition for motor clinical severity [14]. MDS, GMFM, genotype and ages at GMFM and MRI acquisitions are summarized in Table 1. At the first MRI, median age was 8.6 years (2.3 to 43.5). Quantitative DTI measures were obtained in 32 out of 35 patients at median age 8.6 years (2.3 to 41.0). Tractography was carried on in 24 out of 35 patients at median age 13.6 years (2.3–43.5). For some patients, we could not analyze quantitative DTI (n=3) or tractography (n=11) maps due to corrupted data after MRI acquisition. Data were considered corrupted when DTI and tractography software could not read DICOM series because of artefacts due to subject motion. For eight patients, the quantitative DTI and tractography were analyzed on two different MRI scans at different ages in order to obtain non-corrupted data.

Diffusion imaging and parameters

Images were acquired on a 1.5-Tesla Siemens MR scanner under sedation. DTI acquisition used a single-shot echo-planar dual spin-echo sequence: repetition time (TR) =3600ms, echo time (TE) =85ms, 90° flip angle; six diffusion gradients; b0=600 s/mm²; 35 joined axial slices; 256×256 matrix; voxel size=2×2×6mm³; total acquisition time=6 min. T1-weighted, T2-weighted and fluid-attenuated inversion recovery (FLAIR) sequences were added for anatomical guidance (see Supplementary Material 1 for acquisition details).
The DTI parameters fractional anisotropy (FA), mean diffusivity (MD), axial diffusivity (AD) and radial diffusivity (RD) were computed with DTI studio (Jiang and Mori, Johns Hopkins University, cmrm.med.jhmi.edu or godzilla.kennedykrieger.org) using a multivariate linear fitting algorithm for diffusion tensor calculation [15]. We explored 26 regions of interest (ROI), 17 in supratentorial and infratentorial WM and 9 in cortical and deep grey matter (GM) (listed in Supplementary Material 2), after automatic correction of eddy current distortion and automatic segmentation of ROI on the diffusion data based on Harvard-Oxford cortical and subcortical structural atlases (FSL, FMRIB Software Library, fsl.fmrib.ox.ac.uk).

**Tractograms**

Tractography was computed after correction of eddy current distortion, co-registration (mutual information algorithm) and anterior commissure - posterior commissure alignment. Right and left tractograms were generated by a trained neurologist (CS), using fiber assignment by continuous tracking (FACT) [16] and tensor deflection (TEND) [17] methods implemented within iPlan Stereotaxy 3.0 software (BrainLab, Feldkirchen, Germany). FA threshold was set at 0.20 with minimum fiber length tracking of 40 mm. For each tractogram, volume and number of fibers (NOF) were computed and volume/ total intracranial volume (TIV) and NOF/TIV ratios were calculated. TIV was preferred over brain volume for normalization in order to consider brain growth in children and progressive atrophy in adults. The tractograms were interpreted at individual level and visually compared against two control subjects (aged of 8 and 41 years) by two experienced neurologists (CS and JYL) using the following morphologic criteria of fibers bundles: (1) size: large, small, lacking; (2) direction of main-stream fibers (DMSF): anterior-posterior (longitudinal), lateral-medial (transversal), superior-inferior (vertical); (3) connectivity, irrespective of direction, e.g. fugal or petal: extratelencephalic projections (ETP), commissural fibers (CF) and intrahemispheric connections (IHC).

**Statistical analyses**

Statistical analysis was performed using Stata 13 software (Stata-Corp, College Station, TX). The tests were two-sided, with a type-I error set at α=0.05. Quantitative data, expressed as mean and standard deviation or median and min–max ages according to statistical distribution, were compared between three independent groups (PMD0-1, PMD2, and PMD3-4) using a Kruskal–Wallis test followed by the appropriate post-hoc Dunn’s test. ANCOVA was performed with age as the adjustment covariate. The assumption of the normality of residuals was also studied using the Shapiro–Wilk test. Data exhibiting non-normal distributions, a log-transformation was proposed to achieve normality of several variables and to allow the correct use of these analyses. We chose to
report all the individual $p$-values without doing any mathematical correction for distinct tests comparing groups [18]. A focus was given to the magnitude of improvement and to the clinical relevance [19]. Results were expressed as effect-sizes and 95% confidence intervals, and can be compared to the effect-size bounds proposed by Cohen [20], i.e. small ($\pm 0.2$), medium ($\pm 0.5$) and large ($\pm 0.8$, “grossly perceptible and therefore large”). Finally, a nonparametric Spearman’s correlation test, with Sidak’s type-I error correction, was used to compare individual quantitative DTI parameters and GMFM.

**Results**

**Patient severity**

The best motor developmental score (MDS) of the 32 patients analyzed with quantitative DTI were classified as follows: PMD0-1 (n=8), PMD2 (n=14) and PMD3-4 (n=10), with median ages of 8.6 years (3.5–17.4), 6.0 years (2.3–19.1) and 15.4 years (5.5–41.0), respectively. Tractography analysis were possible in 3 PMD0-1 (median ages of 8.5 years (3.0–17.4)), 12 PMD2 (median ages 7.7 years (2.3–43.5)) and 9 PMD3-4 (median age 17.4 years (10.8–37.0)) patients.

The level of the motor handicap at the time of the MRI acquisition was evaluated in the 32 patients using the gross motor function measure (GMFM). Comparison of mean GMFM adjusted to the age between the three MDS groups showed that GMFM was higher in PMD3-4 than PMD0-1 and PMD2. It was also higher in PMD2 than PMD0-1 but not significantly (Fig. 1).

**Quantitative diffusion parameters**

Results of FA, MD, AD, RD in 17 ROI of WM and 9 ROI of GM are shown in Figure 2 (see Supplementary Table 1 for values and Supplementary Fig.1 for effect-sizes and 95% confidence intervals). We found that FA was significantly lower in PMD2 than PMD3-4 in middle cerebellar peduncle, in PMD0-1 than PMD2 in middle temporal gyrus, right caudate nucleus and right thalamus, and in PMD0-1 than PMD3-4 in postcentral gyrus. MD was significantly higher in PMD0-1 than PMD2 in superior cerebellar peduncle, frontal pole and left caudate nucleus, and in PMD0-1 than PMD3-4 in superior cerebellar peduncle. AD was significantly higher in PMD0-1 than PMD2 in superior cerebellar peduncle, left posterior and right superior corona radiata, frontal pole and left caudate nucleus, and in PMD0-1 than PMD3-4 in left superior cerebellar peduncle and right superior corona radiata. RD was significantly higher in PMD0-1 than PMD2 in superior cerebellar peduncle, frontal pole, middle temporal gyrus, caudate nuclei and thalami, and in PMD0-1 than PMD3-4 in right superior cerebellar peduncle and right caudate nucleus.
Motor abilities, evaluated by the GMFM performed at the time of MRI, showed a relatively strong negative linear correlation with MD ($p<0.043$) and AD ($p<0.036$) in the superior cerebellar peduncles. Other possible strong linear relations were observed with MD, AD or RD in anterior corona radiata but were not statistically significant (Fig. 3; Supplementary Table 2).

**Tractograms**

Volume/TIV and NOF/TIV ratios were higher in PMD 3-4 than in other groups ($p<0.02$) after adjustment to age, considering an older age of PMD 3-4 patients at MRI acquisition (Fig. 4, Supplementary Table 3). ETP appeared relatively preserved in all groups. CF were variably individualized in all groups, ranging from large fibers (n=9) to small fibers (n=8) or a lack of connectivity (n=7). IHC size was clinical severity-dependent, being clearly smaller (n=8) or lacking connectivity (n=4) in PMD 2 and PMD 0-1 patients. DMSF of IHC was analyzable in 9 out of 9 PMD 3-4 patients and 10 out of 12 PMD 2 patients, and analysis showed major tracking of longitudinal fibers in 18 out of 19 patients whereas one patient had major transversal fibers (Fig. 5A; Supplementary Fig. 2 and 3). There were age-dependent differences in the PMD 2 group, with smaller CF and IHC in the youngest and oldest patients while patients in late childhood had the largest projections (Fig. 5B; Supplementary Fig. 2 and 3).

**Discussion**

Here we analyzed quantitative DTI measures and tractography patterns from acquisitions performed at the same medical center in a large cohort of PLP1-mutated male patients in order to investigate the ability of these technics to distinguish severity groups. Our 35 patients were representative of the broad clinical spectrum observed in PLP1-related disorders: 46% classical PMD 2 forms, 26% severe PMD 0-1 forms, and 28% mild PMD 3-SPG2 forms. For clinical classification, we used the MDS previously demonstrated to correlate to genotype [3]. To correlate motor skills at the time of MRI, we used GMFM [14], a validated clinical score in cerebral palsy that is widely used in the quantification of motor abilities in leukodystrophic patients. Comparison of mean GMFM between MDS groups argued as in CP for a correlation between the level of the best motor development always observed before 5 years of age and the severity of the motor disabilities later in life. This study shows that quantitative DTI parameters in supratentorial WM are insufficient to distinguish clinical severity in PLP1-mutated patients, whereas increased isotropic diffusion in cortical and deep GM and in infratentorial WM was clinical severity-linked. We also found clinical severity-dependent differences on quantitative and descriptive tractography.
Laukka et al. previously identified increased MD and RD and decreased FA in supratentorial WM regions in 12 PMD patients compared to controls as expected in hypomyelinating brains [13]. Steenweg et al. also found that RD in supratentorial WM was inversely correlated with motor handicap in a cohort of 28 patients with various hypomyelinating disorders [21]. Here, comparison of the MDS groups of our PLP1-mutated patients failed to differentiate clinical severity using quantitative DTI parameters in supratentorial WM. A previous study in the same cohort of patients was unable to classify clinical severity using a myelination score on conventional MRI [9]. Moreover, motor abilities at the time of MRI using GMFM did not significantly decrease with increasing diffusion in supratentorial WM. These results confirmed that increased diffusion and decreased myelination score on the supratentorial WM demonstrate the myelin defect but may not serve to classify the clinical severity observed in PLP1-related disorders. Factors that may have limited the ability of this study to identify differences include the small sample sizes in each group, the broad age-ranges in the patients, or the algorithms and tools used in the quantitative analyses. Other factors could be an offset between microarchitectural changes and clinical features.

Progression in both motor acquisitions and myelination are observed in the first decade of life but followed by motor and cognitive decline and progressive cerebral atrophy [9], so microarchitectural changes in supratentorial WM may precede clinical changes, creating a time gap between diffusion data and clinical data.

In infratentorial WM, clinical severity of the MDS groups and motor abilities at the time of MRI are in accordance with the DTI parameter values, except FA. This result was surprising, as infratentorial WM is usually well myelinated on structural MRI in PLP1-related disorders regardless of clinical severity [19]. However, Laukka et al. also found changes in MD and RD in cerebellar peduncle of PMD patients compared to controls [13]. Moreover, infratentorial abnormalities had been described in a DTI study by Hassen et al. that found MD, AD and RD variations in spinal cord of a SPG2 mouse model [22]. These results suggest that PLP1-mutated patients present subtle changes in infratentorial WM microarchitecture. This idea is consistent with (i) the early and predominant signs of cerebellar dysfunctions observed in PMD patients, (ii) the correlation we had previously found between clinical severity and cerebellar atrophy [9], and (iii) neuropathological analysis of PMD patients with cerebellar abnormalities [23]. In this study, we found higher AD and RD in the most severe patients. Laukka et al. also found changes in AD, which is supposed to reflect axonal integrity, in only the most severe patients, whereas RD, which may reflect myelin integrity, was increased in all forms [13].
In cortical and deep GM, clinical severity of MDS groups increased with the increase of diffusion (MD, AD and RD) in several regions. Moreover, diffusion was more isotropic (decreased FA) in GM in most severe patients. To date in PLP1-related disorders, there are no reports of DTI analysis of GM but Laukka et al. and our team already showed a clear cerebral atrophy including GM compartment and correlating with clinical severity [9,13]. Hassen et al. also found MD increase in GM of spinal cord in a SPG2 mouse model [22]. Increased diffusion in cerebral GM has been reported in multiple sclerosis, where it appears to correlate to motor and cognitive decline [24,25], and in an animal model of genetic demyelinating disorder [26]. However, contrary to our patients, isotropic diffusion decreased (increased FA) in the cortex in multiple sclerosis as a probably result of local activation and proliferation of microglia characterized by the loss of ramifications and acquiring a more anisotropic, bipolar oriented structure as observed in active acute and chronic cortical lesions [25]. In PLP1-related disorders, it may corroborate with (i) the neuron loss, demyelination and axonal damage reported on neuropathological analysis of PLP1-mutated animal models [27] and PMD/SPG2 patients [23], (ii) the absence of local microglial activation and (iii) the neuronal PLP1 expression found in mouse and humans [28,29]. These results underline that GM is clearly involved in the severity of PLP1-related disorders.

Tractography was possible in PLP1-related disorders, including severe forms. Quantitative parameters including tractogram volumes and number of fibers, which have already been used in previous studies [30,31], permitted to distinguish the mildest forms from other forms. Descriptive analyses of fiber bundles showed a preservation of ETP and an altered organization of intra- and interhemispheric connections in all patients, mostly with a loss of vertical and transversal connectivity and an apparent over-representation of longitudinal fibers. This suggests that tractography assessment is easy in mono-directional and partially myelinated fibers but remains difficult in multi-directional and less myelinated fibers in hypomyelinating disorders. Descriptive analyses also revealed an age-dependent evolution of the tractograms in the largest classical–intermediate group. For the youngest patients, the increasing size of tractograms may correspond to different stages of myelination. While healthy 2-year-old children have already achieved myelination except in subcortical fibers, severe supratentorial hypomyelination is the main feature in PLP1-related disorders [10] where myelination can partially improve until 12 years of age [9]. For adult patients presenting severe cerebral atrophy on conventional MRI, the decreased size of tractograms may reflect WM atrophy or loss of connectivity. Laukka et al., argued prominent WM atrophy in PMD/SPG2 patients [11], whereas we observed parallel WM/GM atrophy over time on conventional MRI [9]. Taken together, the combination of patients’ clinical evolution, progressive global atrophy,
GM microstructure changes on DTI and progressive WM fiber disappearance on tractography converge to confirm that \textit{PLP1}-related disorders are global CNS diseases affecting both WM and GM.

This study has several limitations. First, recent studies suggested the potential for seemingly minor alterations in data acquisition, in particular the b-value and number of diffusion gradient directions, to impact the DTI parameters [32]. Moreover, using multiple b-values such as recent technique called neurite orientation dispersion and density imaging (NODDI) could provide additional information on tissue compartments [33]. Further studies should consider these parameters. Second, with only 6 gradient directions, tractography can be limited to only major fiber tracts and can underestimate the connectivity. Our goal to differentiate severity in \textit{PLP1}-related disorders was achieved in this protocol but it would be interesting to increase the number of directions to study weak connections. Finally, even if we took into account the age of the subjects as a covariate, there is a difference of age between the patients’ groups that could generate a bias of interpretation. We did not report intra-individual evolution of DTI parameters and tractography. This would be a different study that could help to strengthen our results.

In conclusion, this study shows for the first time that tractography can be a useful and non-invasive technique in investigations and follow-up of hypomyelinating disorders. Our findings also bring further evidence of physiopathological mechanisms in \textit{PLP1}-related disorders that likely involve both grey and white matter and infratentorial structures. These findings must be confirmed in further studies and may be important in helping to develop new therapeutic strategies targeting not only oligodendrocytes but also axons and neurons.
Acknowledgements
The authors thank the members of the Leukofrance network who send us the patients (MA Barthez-Carpentier, F Boidin, P Burkart, S Ceylaner, B Chabrol, D Chaigne, G Cioni, B Echenne, M Elleder, P Evrard, M Garcia Silva, K Kluger, J M Lopez-Terradas, J Motte, A Munnich, D Nicholls, N Philip, G Ponsot, T Reckert, D Rodriguez, C Rousselle, G Sébire, H Steinbock, M Troncoso, H Van Esche, and L Van Maldergem) and Dr JP Renou for precious help with the DTI analyses.

Funding
This study received funding from the ELA foundation, the FP7-RD Connect, the national PHRC [hospital-based clinical research program] for Clermont-Ferrand university hospital, and the Région Auvergne regional council.
JJL, CS, AS, JC, NS report a team non-financial support from BrainLab. The other authors declare no competing interests.

List of Supplementary Material
- Supplementary Material 1. Details of acquisition of T1-weighted, T2-weighted and FLAIR sequences.
- Supplementary Material 2. List of ROI studied for quantitative DTI.
- Supplementary Table 1. Results of statistical analyses for quantitative DTI parameters in white and grey matter.
- Supplementary Table 2. Results of statistical correlation between GMFM and quantitative DTI parameters in white and grey matter.
- Supplementary Table 3. Results of statistical analysis for quantitative parameters of left and right tractograms in each group of patients.
- Supplementary Figure 1. Additional results of statistical analyses for the different biomarkers: representation of effect sizes.
- Supplementary Figure 2. Superior views of both hemisphere tractograms of different severity groups and ages in PMD/SPG2 patients.
- Supplementary Figure 3. Lateral left views of right hemisphere tractogram of different severity groups and ages in PMD/SPG2 patients.
References


### Table 1. List of patients and families with PMD/SPG2

<table>
<thead>
<tr>
<th>Patient</th>
<th>Family</th>
<th>MDS</th>
<th>Age at DTI acquisition +/- GMFM (years)</th>
<th>GMFM88 at MRI acquisition</th>
<th>PLP1 mutation</th>
<th>Family genetic number*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>2</td>
<td>2.3</td>
<td>29.3</td>
<td>duplication</td>
<td>1910</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>0</td>
<td>3.0</td>
<td>NA</td>
<td>c.454-1G&gt;A</td>
<td>767</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>1</td>
<td>3.5</td>
<td>36.8</td>
<td>c.327delC</td>
<td>1171</td>
</tr>
<tr>
<td>4</td>
<td>4</td>
<td>2</td>
<td>3.8</td>
<td>25.3</td>
<td>duplication</td>
<td>NA</td>
</tr>
<tr>
<td>5</td>
<td>5</td>
<td>2</td>
<td>4.3</td>
<td>34.8</td>
<td>duplication</td>
<td>1251</td>
</tr>
<tr>
<td>6</td>
<td>6</td>
<td>2</td>
<td>4.4 / 6.8</td>
<td>26 / 36.8</td>
<td>duplication</td>
<td>1059</td>
</tr>
<tr>
<td>7</td>
<td>7</td>
<td>2</td>
<td>4.5 / 7.5</td>
<td>NA / 39.5</td>
<td>c.659G&gt;A</td>
<td>759</td>
</tr>
<tr>
<td>8</td>
<td>8</td>
<td>2</td>
<td>4.7</td>
<td>NA</td>
<td>duplication</td>
<td>1201</td>
</tr>
<tr>
<td>9</td>
<td>9</td>
<td>2</td>
<td>5.1</td>
<td>38.4</td>
<td>duplication</td>
<td>1026</td>
</tr>
<tr>
<td>10</td>
<td>10</td>
<td>3</td>
<td>5.5</td>
<td>39</td>
<td>duplication</td>
<td>1114</td>
</tr>
<tr>
<td>11</td>
<td>11</td>
<td>1</td>
<td>5.7</td>
<td>28.7</td>
<td>duplication</td>
<td>968</td>
</tr>
<tr>
<td>12</td>
<td>12</td>
<td>2</td>
<td>5.7</td>
<td>26.7</td>
<td>duplication</td>
<td>887</td>
</tr>
<tr>
<td>13</td>
<td>13</td>
<td>2</td>
<td>6.2</td>
<td>47.1</td>
<td>c.125delG</td>
<td>1038</td>
</tr>
<tr>
<td>14</td>
<td>14</td>
<td>3</td>
<td>8.3 / 11.5</td>
<td>44.2 / NA</td>
<td>duplication</td>
<td>407</td>
</tr>
<tr>
<td>15</td>
<td>15</td>
<td>0</td>
<td>8.5</td>
<td>20.5</td>
<td>duplication</td>
<td>545</td>
</tr>
<tr>
<td>16</td>
<td>16</td>
<td>1</td>
<td>8.6</td>
<td>17</td>
<td>duplication</td>
<td>1776</td>
</tr>
<tr>
<td>17</td>
<td>17</td>
<td>0</td>
<td>8.6</td>
<td>NA</td>
<td>c.670C&gt;T</td>
<td>766</td>
</tr>
<tr>
<td>18</td>
<td>18</td>
<td>2</td>
<td>8.6</td>
<td>41.4</td>
<td>duplication</td>
<td>418</td>
</tr>
<tr>
<td>19</td>
<td>19</td>
<td>3</td>
<td>10.8 / 13.3</td>
<td>NA / 45</td>
<td>c.173A&gt;G</td>
<td>106</td>
</tr>
<tr>
<td>20</td>
<td>20</td>
<td>1</td>
<td>12.3</td>
<td>NA</td>
<td>duplication</td>
<td>440</td>
</tr>
<tr>
<td>21</td>
<td>21</td>
<td>3</td>
<td>13.5</td>
<td>NA</td>
<td>duplication</td>
<td>46</td>
</tr>
<tr>
<td>22</td>
<td>22</td>
<td>2</td>
<td>13.6</td>
<td>46.9</td>
<td>duplication</td>
<td>154</td>
</tr>
<tr>
<td>23</td>
<td>23</td>
<td>3</td>
<td>14.5 / 17.4</td>
<td>49.9 / 30.5</td>
<td>c.307delG</td>
<td>905</td>
</tr>
<tr>
<td>24</td>
<td>24</td>
<td>2</td>
<td>14.8</td>
<td>40</td>
<td>duplication</td>
<td>61</td>
</tr>
<tr>
<td>25</td>
<td>25</td>
<td>1</td>
<td>15.0</td>
<td>NA</td>
<td>duplication</td>
<td>150</td>
</tr>
<tr>
<td>26</td>
<td>26</td>
<td>4</td>
<td>16.2</td>
<td>56.6</td>
<td>c.548C&gt;A</td>
<td>41</td>
</tr>
<tr>
<td>27</td>
<td>27</td>
<td>2</td>
<td>16.8 / 19.1</td>
<td>0 / 0</td>
<td>c.608A&gt;G</td>
<td>37</td>
</tr>
<tr>
<td>28</td>
<td>28</td>
<td>2</td>
<td>17.0</td>
<td>24</td>
<td>duplication</td>
<td>28</td>
</tr>
<tr>
<td>29</td>
<td>29</td>
<td>3</td>
<td>17.2 / 20.1</td>
<td>44.4 / 32.3</td>
<td>c.307delG</td>
<td>905</td>
</tr>
<tr>
<td>30</td>
<td>30</td>
<td>0</td>
<td>17.4</td>
<td>NA</td>
<td>c.454-1G&gt;T</td>
<td>26</td>
</tr>
<tr>
<td>31</td>
<td>31</td>
<td>3</td>
<td>19.0</td>
<td>24</td>
<td>c.646C&gt;G</td>
<td>53</td>
</tr>
<tr>
<td>32</td>
<td>32</td>
<td>1</td>
<td>21.8</td>
<td>NA</td>
<td>c.646C&gt;T</td>
<td>24</td>
</tr>
<tr>
<td>33</td>
<td>33</td>
<td>3</td>
<td>32.8</td>
<td>NA</td>
<td>c.173A&gt;G</td>
<td>106</td>
</tr>
<tr>
<td>34</td>
<td>34</td>
<td>4</td>
<td>37.0 / 41.0</td>
<td>31.8</td>
<td>c.801G&gt;T</td>
<td>1</td>
</tr>
<tr>
<td>35</td>
<td>35</td>
<td>7</td>
<td>43.5</td>
<td>NA</td>
<td>c.659G&gt;A</td>
<td>759</td>
</tr>
</tbody>
</table>

MDS: motor developmental score, age (years) at DTI acquisition and GMFM scoring, GMFM: gross motor function measure realized at the time of MRI acquisition, and genotype (at the DNA level).

*Family genetic number corresponds to our laboratory family number and refers to previous publication (Cailloux et al. 2000). NA : not available.
**Figure 1.** Comparison of mean GMFM between the three MDS groups.
Figure 2. Comparison of quantitative DTI parameters between the three MDS groups in white matter, subcortical white matter and grey matter.
Figure 3. Coefficients of correlation between GMFM and quantitative DTI parameters in white and grey matter.

Figure 4. Comparison of quantitative tractography parameters between the three MDS groups.
Figure 5. Examples of frontal views of both hemisphere tractograms and results of morphologic analyses of tractographies in different severity groups and ages in PMD/SPG2 patients.
Figure legends

**Figure 1.** Comparison of mean GMFM between the three MDS groups. **Y-axis:** GMFM (Gross Motor Function Measure) includes 88 items (best motor score = 88) at the time of MRI acquisition. **X-axis:** MDS (Motor developmental score) corresponds to the best motor function achieved between the ages of 2 and 5 years. Thirty-two patients are divided in three groups of MDS: PMD0-1 (n=8), PMD2 (n=14), PMD3-4 (n=10). Results are given using ANCOVA with age as the adjustment covariate. * indicates a significant difference (p-value <0.05). p = p-value. Standard deviations are given as an error bar.

**Figure 2.** Comparison of quantitative DTI parameters between the three MDS groups in white matter, subcortical white matter and grey matter. Successive results for fractional anisotropy (FA), mean diffusivity (MD), axial diffusivity (AD) and radial diffusivity (RD). Thirty-two patients are divided in three groups of MDS: PMD0-1 (n=8), PMD2 (n=14), PMD3-4 (n=10). MD, AD and RD values are expressed as $10^{-3}$ values. Results are expressed as means. Standard deviations are given as an error bar. * for $p$-value <0.05, $p$-values were computed using ANCOVA with age as the adjustment covariate. ALIC: anterior limb of the internal capsule, PLIC: posterior limb of the internal capsule, R: right, L: left, CC: corpus callosum, Mid: middle, Ant: anterior, Post: posterior, Sup: superior, Ped: peduncle, Cereb: cerebellar.

**Figure 3.** Coefficients of correlation between GMFM and quantitative DTI parameters in white matter, subcortical white matter and grey matter. Using a nonparametric Spearman’s correlation test, with Sidak’s type-I error correction, values of coefficients of correlation between -1 and 1 are represented graphically with a color-coded heatmap in each region of interest and for each DTI parameters. GMFM: gross motor function measure, FA: fractional anisotropy, MD: mean diffusivity, AD: axial diffusivity, RD: radial diffusivity, ALIC: anterior limb of internal capsule, PLIC: posterior limb of internal capsule, R: right, L: left, CC: corpus callosum, Mid: middle, Ant: anterior, Post: posterior, Sup: superior, Ped: peduncle, Cereb: cerebellar.

**Figure 4.** Comparison of quantitative tractography parameters between the three MDS groups. Successive results for total intracranial volume (TIV), right and left tractogram volume / TIV and right and left tractogram number of fibers / TIV. Twenty-four patients whom tractography was available are divided in three groups of MDS: PMD0-1 (n=3), PMD2 (n=12), PMD3-4 (n=9). Results are expressed as means. Standard deviations are given as an error bar. Total intracranial.
volume (TIV) and tractogram volume are expressed in mm$^3$. * for p-value <0.05. p-values were computed using ANCOVA with age as the adjustment covariate.

**Figure 5.** Examples of frontal views of both hemisphere tractograms and morphological analyses of tractographies of different MDS groups and ages in PMD/SPG2 patients. Tractograms were realized for two regions of interest corresponding to the right/left hemispheres. Colors correspond to the direction of main-stream fibers (DMSF): blue = vertical direction, green = longitudinal direction, red = transversal direction. For clinical comparison, Gross Motor Functional Measure (GMFM) at the time of MRI is given under each tractogram when available. A. Examples of tractograms for each group of severity and table of the results of morphological analyses in 24 patients whom tractography was available: PMD0-1 (n=3), PMD2 (n=12) and PMD3-4 (n=9). Extratelencephalic projections (ETP) were relatively preserved in all groups, but were smaller than in controls. Compared to controls (a,e), commissural fibers (CF) and intrahemispheric connections (IHC) were smaller or lacking in all groups: (1) the most severe PMD0-1 (b,f), (2) the classical PMD2 (c,g) and the mildest PMD3-4 patients (d,h). DMSF in IHC was marked by longitudinal (or rare transversal) over representation. B. Each image corresponds to tractograms of PMD2 patients of different ages. These results show that (1) the youngest patients (a,b) had less-developed ETP, CF and IHC than patients at the late infancy (c,d) and adolescent patients (e,f); (2) the oldest patients (g,h) had smaller ETP, CF and IHC than adolescent patients (e,f).