Linking white matter and deep gray matter alterations in premanifest Huntington disease

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A B S T R A C T

Huntington disease (HD) is a fatal progressive neurodegenerative disorder for which only symptomatic treatment is available. A better understanding of the pathology, and identification of biomarkers will facilitate the development of disease-modifying treatments. HD is potentially a good model of a neurodegenerative disease for development of biomarkers because it is an autosomal-dominant disease with complete penetrance, caused by a single gene mutation, in which the neurodegenerative process can be assessed many years before onset of signs and symptoms of manifest disease. Previous MRI studies have detected abnormalities in gray and white matter starting in premanifest stages. However, the understanding of how these abnormalities are related, both in time and space, is still incomplete. In this study, we combined deep gray matter shape diffeomorphometry and white matter DTI analysis in order to provide a better mapping of pathology in the deep gray matter and subcortical white matter in premanifest HD. We used 296 MRI scans from the PREDICT-HD database. Atrophy in the deep gray matter, thalamus, hippocampus, and nucleus accumbens was analyzed by surface based morphometry, and while white matter abnormalities were analyzed in (i) regions of interest surrounding these structures, using (ii) tractography-based analysis, and using (iii) whole brain atlas-based analysis. We detected atrophy in the deep gray matter, particularly in putamen, from early premanifest stages. The atrophy was greater both in extent and effect size in cases with longer exposure to the effects of the CAG expansion mutation (as assessed by greater CAP-scores), and preceded detectible abnormalities in the white matter. Near the predicted onset of manifest HD, the MD increase was widespread, with highest indices in the deep and posterior white matter. This type of in-vivo macroscopic mapping of HD brain abnormalities can potentially indicate when and where therapeutics could be targeted to delay the onset or slow the disease progression.

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1. Introduction

Huntington’s disease (HD) is a devastating progressive neurodegenerative disorder that affects multiple domains, including motor, cognitive, and emotional, leading to incapacity for activities of daily living and eventually, to death (Huntington, 1872; Folstein, 1991; Ross et al., 2014). It is caused by CAG repeat expansion in the gene Huntington (HHTT), leading to polyglutamine expansion in the protein huntingtin (The Huntington’s Disease Collaborative Research Group, 1993). Starting at the threshold of 36 CAGs, longer expansions lead to earlier ages of motor onset of HD. Diagnosis of “motor onset” of “manifest HD” is currently made in someone at risk, or tested genetically positive for the CAG expansion, on the basis of the clinical history and standardized Unified HD Rating Scale (UHDRS) motor exam yielding a clinical impression with “99% confidence” of the presence of HD in the context of the unequivocal presence of an otherwise unexplained extrapyramidal movement disorder (The Huntington’s Disease Collaborative Research Group, 1993; The Huntington’s Disease
The course of HD can be divided into ‘premanifest’ and ‘manifest’ periods (Zhang et al., 2011; Dorsey et al., 2013; Tabrizi et al., 2013; Paulsen et al., 2014a, 2014b; Ross et al., 2014). The premanifest period can be further subdivided (Reilmann et al., 2014; Ross et al., 2014), and we use terminology from these references. First is a period when individuals are not distinguishable clinically (either by subjective symptoms or objective findings on exam) from controls (“presymptomatic”), usually up to 10–15 years before “motor onset.” Individuals may then enter the “prodomal” period, which is characterized by subtle motor, cognitive and behavioral changes, but not sufficient for a diagnosis of motor onset. Once motor and cognitive signs and symptoms begin, they progress inexorably over the course of the illness, which—with the exception of late-onset cases, who may die of other causes—is uniformly fatal (Ross et al., 1997; Dorsey et al., 2013).

A useful index of the length and severity of the individual’s exposure to the effects of the mutant HTT gene is achieved by calculating a score based on the individual’s age multiplied by (CAG – 1), where L is a constant near the threshold of CAG repeat expansions for disease (Penney et al., 1997). This is termed the CAG–Age–Product (or CAP score). This score is useful for comparing data from cohorts of patients with a range of ages and CAG repeat lengths (Zhang et al., 2011; Ross et al., 2014).

There is growing consensus that intervention and treatment in HD should occur at the earliest stage possible. The development and testing of neuroprotective treatments will be facilitated by the identification of biomarkers in the manifest HD period and, optimally, premanifest stages. Neuroprotective treatments could include administration of gene silencing or other reagents directly into localized regions of the brain (Huntington’s Disease Collaborative Research Group et al., 2014; Kay et al., 2014; Ross et al., 2014). Because of the inverse relationship between the length of the CAG repeat expansion and age of onset, HD is a model disease that may provide proof of concept for neurodegenerative disease therapeutic modification. It also involves increasingly well-understood natural history, as demonstrated by large multicenter studies such as TRACK-HD (Tabrizi et al., 2009, 2012, 2013) and PREDICT-HD (Paulsen et al., 2006, 2008, 2014a, 2014b). These studies have identified white abnormalities in pre-manifest stage, in addition to the well-known atrophy of striatum and other subcortical gray matter structures.

Structural imaging from PREDICT, TRACK-HD, and other studies revealed striatal atrophy starting as early as 15 years before the onset, and continuing throughout the pre-manifest periods (Aylward et al., 1997, 2000, 2003, 2011, 2012; Paulsen et al., 2010, 2014a, 2014b; Tabrizi et al., 2012, 2013). Extent of striatal atrophy adds predictive power for motor onset beyond age and CAG repeat length alone (Aylward et al., 2012, 2013; Paulsen et al., 2014a, 2014b). In addition, the measurement of shape has enabled the delineation of regional atrophy in the striatum and other subcortical structures in the premanifest period (Younes et al., 2014a, 2014b). In parallel, diffusion tensor imaging (DTI) has revealed abnormalities in architecture and integrity in white matter and subcortical gray matter structures in both premanifest (Reading et al., 2005; Rosas et al., 2006; Koppel et al., 2008; Stoffers et al., 2010) and manifest phases (Douaud et al., 2009; Vandenberge et al., 2009; Della Nave et al., 2010; Bohanna et al., 2011; Delmaire et al., 2013; Sanchez-Gastaneda et al., 2013).

In this study, we combine deep gray matter shape diffeomorphometry and white matter DTI analysis as means of better defining the topography of deep gray and white matter abnormalities in the premanifest stages of HD. Diffeomorphometry and geodesic shape analysis in computational anatomy (Van Camp et al., 2012; Miller et al., 2013, 2014; Younes et al., 2014a, 2014b) provide information about which structure subregions are affected. The association of this metric with DTI parameters in white matter regions through the pre-manifest course may provide additional details about the mapping of brain abnormalities. This could provide information about when, where, and how therapeutics could be administered in order to delay the onset or slow the progression of HD.

2. Methods

2.1. Participants

The data used here are from the PREDICT-HD study, in which subjects at risk for HD, but without motor signs and symptoms sufficient for diagnosis of manifest HD, had previously undergone elective predictive genetic testing. Premanifest and prodromal subjects are those that are found to be gene-expanded (CAG length greater than or equal to 36) but have not yet met traditional motor criteria for a diagnosis of HD. Diagnosis of HD is based on a Diagnostic Confidence Level (DCL) rating of “4” according to the motor assessment section of the Unified HD Rating Scale (UHDRS). Those that were found to negative for the CAG expansion (CAG at or below 30) were enrolled as comparison subjects. There were no subjects in with CAG between 30 and 36. Participants were recruited from 33 sites across the United States, Canada, Europe, and Australia, and had annual study visits consisting of a neurological motor examination, cognitive assessment, brain MRI (biennial), and psychiatric and functional assessment, with blood samples for genetic and biochemical analyses. Informed written consent was obtained from all subjects before participating in this study.

Subjects with CAG expansion were divided into three subgroups: low-CAP, medium-CAP, and high-CAP. Low-CAP individuals are presumably far from the symptomatic onset while the high-CAP individuals are the closest to the onset. The CAP score was computed as $\text{CAP} = \text{age at entry} \times (\text{CAG} - 33.66)$ (Zhang et al., 2011) and presumably indexes the cumulative exposure to the effects of mutant huntingtin. Cutoffs for groups were CAP < 290 (Low), 290 ≤ CAP ≤ 368 (Medium), and CAP > 368 (High).

2.2. MRI

High-resolution structural 3T MRI scans (T1-weighted inversion recovery turboflash (MPRAGE), were used for the analyses of diffeomorphometry. Due to the multicentre and longitudinal nature of the PREDICT study, the image acquisition was heterogeneous, represented by multiple vendors (GE, Phillips, and Siemens) and over 20 slightly different MR acquisition protocols (i.e., due to transmission and receive hardware). The main parameters of the data included in this study varied as follows: TR: 8–8.4, TE: 3.5–3.8, TI: 826–843, matrix: 256 × 256, voxel size: 1 × 1, slice thickness: 0.9–1.2. Similar data have been successfully used in previous publications authored by the Huntington’s Study Group (HSG) PREDICT investigators (Paulsen et al., 2010, 2014a, 2014b; Muralidharan et al., 2014; Kim et al., 2015).

The DTIs were obtained in 3T scanners, 80 = 1000, in axial orientation (matrix = 128 × 128, FOV = 256 × 256, slice thickness = 2 mm). The number of slices, gradients, and repetitions varied according to the center/scanner: 1) Phillips: 33 gradients, 70 slices, 2 repetitions; 2) Siemens (most common): 31 gradients, 70 slices, 2 repetitions; 3) Siemens (minority): 79 gradients, 50 slices, 3 repetitions. The repetitions were concatenated within a single section using DTIPrep (http://www.nitrc.org/projects/dtiprep/) (Oguz et al., 2014), followed by quality control inspection, and then converted to nifti format using DWICONvert. Details about the tensor calculation and quality control are fully described in our previous publications (Magnotta et al., 2012); similar data was used in (Matsui et al., 2015). The six elements of the diffusion tensor, the fractional anisotropy (FA), and the mean diffusivity (MD) were calculated using DTIstudio software (Laboratory of Brain Anatomical MRI and Center for Imaging Science at Johns Hopkins University), and visual quality control was performed to ensure that the values for scalar images were in proper range and that gross anatomical FA values were appropriate. Our previous publications showed that scalar measures such as FA and MD are stable in face of different gradient number and directions, minimizing the effect of different protocols in multicenter studies (Magnotta et al., 2012).
We used scans of 296 individuals enrolled in the PREDICT-HD study. When more than one session was available for an individual, the first visit was used. Table 1 summarizes the demographic information. Twenty-three DTI scans were not included because the original files were missing or incomplete, or because of rejections at the first step of the quality control, in the tensor and scalar calculations. For the shape diffeomorphometry, the sample size for each group varied by as many as five individuals in the comparison for each substructure due to the quality of the resulting triangulated mesh associated to each substructural segmentation forming the shape statistics. The demographic profile of each group was not significantly different among the comparison of shape in different structures and of the DTI.

### 2.3. Surface-based morphometry of subcortical structures

We extend the statistical analyses of diffeomorphometry followed in previous papers (Csernansky et al., 1998, 2000; Ashburner et al., 2003; Thompson et al., 2004; Wang et al., 2007; Qiu et al., 2009, 2010; Miller et al., 2013, 2014; Younes et al., 2013). This procedure has three steps: (i) segmentation of the target structures, (ii) generation of a single template coordinate system from the population of baseline scans, and (iii) mapping of the template onto each of the target segmented structures represented via triangulated meshes.

The first step (segmentation of the target structures) was done using the BRAINS AutoWorkup processing pipeline published in (Pierson et al., 2011). After completion of AutoWorkup, all scans were individually inspected and manually corrected where necessary to ensure correct realignment and coregistration, tissue classification, and accuracy of brain and subcortical structures. The description of this process and its reliability is fully described in (Kim et al., 2014). The resulting preprocessed data included T1, T2, PD, and tissue classification volumetric images where the anterior commissure (AC point) was set to be the center of the image resampled to 1 mm resolution in a 256 × 256 × 256 matrix. Details of each step of MPRAGE pre-processing are found in our previous publication (Young Kim and Johnson, 2013).

Binary images for the six substructures (caudate, globus pallidus, putamen, thalamus, hippocampus, and nucleus accumbens) were evaluated. To generate shape biomarkers indexed to a common coordinate system, we follow the procedure published in Younes et al. (2013) in which all surfaces are meshed and rigidly aligned via rotation and translation, with right subvolumes flipped before alignment to ensure that all structures could be compared. From rigidly aligned volumes, an average template triangulated mesh was generated based on a generative probability model over the entire population in which the observed surfaces are modeled as random deformations of the template (Ma et al., 2010). This template generation procedure solves a maximum-likelihood estimation problem for a statistical model in which the population is described as a random deformation of the template, itself defined as an unknown deformation of a fixed surface called hypertextemplate. This procedure provides a coordinate system that represents the population via surface mapping onto the template. The resulting templates for the caudate, putamen, globus pallidus, thalamus, hippocampus, and nucleus accumbens (Fig. 1) become the coordinate systems, which are referenced for our p-value statistics and FWER (Family-wise Error Rate) calculations. These were computed by running the template generation algorithm on the population of baseline scans and are blind to group labels.

The high-dimensional diffeomorphometry shape statistics are generated indexed to each location of the template by computing the diffeomorphic correspondence between the template and each surface using large deformation diffeomorphic metric mapping (LDDMM) surface registration (Vaillant and Glaunes, 2005). The algorithm computes a smooth invertible mapping of the triangulated surface template \( S_{\text{temp}} \) onto the target surfaces \( S_{\text{target}} \) by minimizing the geodesic transformation energy and the matching cost that is the distance measurement between the mapped template (\( \phi_t \cdot S_{\text{temp}} \)) and the target surface.

\[
E = \delta(\phi_t \cdot S_{\text{temp}}, S_{\text{target}}) + \alpha \int_0^1 \nu_t^2 dt \text{ with } \nu_t = \nu_t(\phi_t)
\]

The matching cost term \( \delta(\phi_t \cdot S_{\text{temp}}, S_{\text{target}}) \) computes a norm between surfaces.

Shape coordinates are then deduced from the optimal diffeomorphism, resulting in one scalar measure per vertex on the template surfaces. For computational efficiency, we sub-discretized these measures by averaging them over small segments computed on the surface template (Younes et al., 2014a, 2014b). These segments are obtained by spectral clustering of the surface, a method that only relies on the surface geometry. This is achieved by computing the first \( k \) eigenvectors of the Laplace–Beltrami operator associated with the surface, where \( k \) is the intended number of segments, associating with each vertex a \( k \)-dimensional vector formed with the values of the eigenvectors evaluated at this point. These vectors are then used in a standard K-means algorithm to provide the \( k \)-desired segments. The number of segments was adjusted so that they cover an area of 150 mm\(^2\) on average, yielding 10 segments on the putamen, 8 on the caudate, 3 on the globus pallidus, 12 on the thalamus, 1 on the nucleus accumbens, and 6 on the hippocampus.

### 2.4. White matter analysis with DTI

The pipeline for the white matter DTI analysis consisted on mapping each subject's brain to a common brain template (JHU-MNI-SS) (Mori et al., 2008), using a sequence of linear transformations and then LDDMM (Miller et al., 2005; Wang et al., 2007; Ceritoglu et al., 2009). The JHU-MNI-SS is a multi-MRI contrast brain extensively segmented to 12 on the thalamus, 1 on the nucleus accumbens, and 6 on the hippocampus.

### Table 1

Demographic information and sample size used in each comparison.

|                      | Controls | Low-CAP | Medium-CAP | High-CAP | Total |
|----------------------|----------|---------|------------|----------|-------|
| **Total**            | 84       | 58      | 76         | 78       | 296   |
| **DTI**              | 79       | 54      | 68         | 72       | 273   |
| **Diffeomorphometry**|          |         |            |          |       |
| Caudate              | 84       | 57      | 76         | 78       | 295   |
| Putamen              | 81       | 57      | 76         | 73       | 287   |
| Globus pallidus      | 82       | 57      | 75         | 77       | 291   |
| Thalamus             | 84       | 58      | 76         | 74       | 292   |
| Hippocampus          | 79       | 56      | 75         | 73       | 283   |
| Nc. accumbens        | 84       | 58      | 76         | 78       | 296   |
| **Age**              | 46.3 ± 11.1 | 35.1 ± 9.9 | 42.8 ± 9.3 | 47.9 ± 9.8 |   |
| **Gender**           | 53F/31M  | 17F/41M | 23F/53M    | 29F/49M  |       |
and ROIEditor (Laboratory of Brain Anatomical MRI and Center for Imaging Science at Johns Hopkins University).

In order to identify the relationship between deep gray matter and deep white matter in HD, we used three approaches: A) the analysis of regions of interest (ROIs) in the white matter adjacent to the deep gray matter; B) the analysis of white matter pathways between the deep gray matter and cortex, using dynamic programming (DP); C) the analysis of the total white matter, regionally parcellated, using an atlas-based approach (ABA) (Fig. 2).

2.4.1. Analysis of the white matter adjacent to the deep gray matter

Each subregion defined in the surface of caudate, putamen, globus pallidus, and thalamus for the gray matter shape analysis (Fig. 2B, left panel) was expanded radially by 5 voxels (in a 1mm³ image resolution) through the adjacent white matter. Therefore, each segment in the shape analysis has the corresponding "white matter neighbor," except where the surfaces are adjacent to the ventricles and in the putamen–globus pallidum interface.

2.4.2. Analysis of the white matter pathways between the striatum and the cortex

White matter pathways were traced using DP between each of the surface segments in striatum used for diffeomorphometry and the 38 cortical parcels in each hemisphere. The cortical parcels were previously defined in the template, as mentioned above. The DP algorithm applied is detailed in our previous paper (Li et al., 2014). Our choice of a probabilistic path-generation algorithm aims to minimize the false negatives that simple line propagation methods would generate, given that our seeds are located in the gray matter. Since high-angular resolution images are not available, DP is the natural option for tracing. Using a "not" operation we removed connections crossing a third seed or the ventricles. This minimized the double sampling of pathways and excluded implausible pathways. The streamlines traced in the template were eventually converted to regions of interest.

It is worth noting that algorithms such as DP trace the "shortest path" between chosen initial and terminal points that globally minimize a sequentially additive energy constraint defined by the tensor and, as happens to any other DTI-based tract-tracing method, does not necessarily reflect the existence of a biological connection. Therefore, they have to be interpreted as a macroscopic reconstruction of the white matter architecture or a region-growing tool, which can cluster anatomically related pixels based on DTI data, rather than a tool with which to investigate connectivity based on a cellular level structure.

2.4.3. Regional analysis of the whole white matter

Using our pre-segmented template and accurate image mapping, we automatically parceled the whole white matter of each individual into 98 ROIs. The segregation of cortex and peripheral white matter was threshold by FA > 0.2. This "atlas-based" analysis carries information about the structural anatomy, since each parcel is defined based on previous anatomical knowledge.

2.5. Group based linear effects statistical analysis

We have performed statistical analyses on each of the subcortical structures, making separate comparisons of the shape diffeomorphometry markers corresponding to the degree of atrophy relative to the template between the groups. We modeled the shape diffeomorphometry markers via linear effects. The analysis includes age, gender and log intracranial volumes as covariates, and computes statistics at each segment of the triangulated template surface returning p-values corrected for multiple comparisons using permutation testing.

The group analysis quantifies a mixed linear effects model of atrophy between the groups including controls and the low, medium and high-
CAP score defined groups, and is given for each vertex \( v \), scan \( j \) and subject \( s \). For each control and patient subject, left and right structures are registered to the template, resulting in the computation of a normalized deformation marker \( J_v(s) \) that measures the amount of expansion/atrophy at vertex \( v \) of the template surface in registering it to subject \( s \). The raw expansion/atrophy measure is defined as the logarithm of the local expansion/reduction in surface area around the vertex, and is interpreted mathematically as a log-jacobian on the template surface. We model the group variables as \( g(s) \) equaling 1 if subject \( s \) belongs to the group (either low, medium or high), and zero otherwise belonging to the controls. Our analysis uses gender (denoted \( d \)) and intracranial volume (denoted \( i \)) as covariates, resulting in the model

\[
J_v(s) = \alpha_v + \beta_v g(s) + \gamma_v d(s) + \delta_v i(s) + \epsilon_v(s)
\]  

(2)

with \( \epsilon_v(s) \) representing our noise model with \( \epsilon_v(s) \) Gaussian distributed with variance \( \sigma^2 \). We test for the null hypothesis with \( H_0^v: \beta_v = 0 \) for all \( v \), while correcting for multiple comparisons.

For volume testing, the logarithm of the volume is used. The p-values of these models are computed using permutation sampling (using random permutations of residuals of the null hypothesis model). The test statistic is the log-likelihood difference between the null hypothesis and the \( H_1^v: \beta_v \neq 0 \) general hypothesis, which is equivalent to computing, for each coordinate \( v \), the logarithm of ratio of the residual variance for the complete \( H_1^v \) hypothesis to the one obtained from the null hypothesis. Denoting this statistic by \( S_v \), the family-wise error rates (FWER) are calculated by evaluating the maximum \( S^* = \max_v S_v \).

To compute p-values, the same statistic \( S^* \) is computed a large number of times with permuted residuals under the null, i.e., taking

\[
J_v^\pi(s) = \alpha_v^\pi + \gamma_v^\pi d(s) + \delta_v^\pi i(s) + \epsilon_v^\pi(s)
\]  

(3)

where \( \pi \) is a random permutation of the subjects and the parameters and residuals are those obtained using maximum likelihood under the null hypothesis. The p-value is given by the fraction of times the values of \( S^* \) computed after permuting the residuals is larger than the value obtained with the true groups. The p-values that were observed via the linear effects modeling of deformation markers of Eq. (2) are provided for each structure including the left and right sides of the brain in the Supplementary Table 1. The volume statistics shown in the third column for the three groups provides p-values for the same linear effects model, also evaluated via permutations, in which \( J \) is replaced by the structure volume, for which no multiple testing correction is required.

For the DTI analysis, after mapping each subject brain scan to the common template, mean values of fractional anisotropy (FA), mean diffusivity (MD), radial diffusivity (RD), and axial diffusivity (AD) were obtained in each region of interest, in each of the three approaches mentioned before. Similarly, differences among groups (controls and each of the premanifest HD) were evaluated with permutation test and corrected for multiple comparisons with FWE. Monotonic linear fitting was used to describe the relationship between shape and DTI parameters versus CAP scores.

3. Results

3.1. Shape diffeomorphometry

The shape diffeomorphometry analysis of subcortical gray matter change demonstrated differences in all three pre-manifest stages versus controls (Fig. 3). Modest differences, both in terms of location (restricted to central segments of putamen) and effect size were detected in the low-CAP group. In the medium-CAP, the differences extended to the whole surface of putamen, caudate, globus pallidus, and nucleus accumbens. In the high-CAP group, differences were also noticed in thalamus.
and hippocampus. The effect size was different in different regions of striatum and other subcortical structures. Especially large atrophy were noted in some regions, such as tail of caudate, showing up to 25% of atrophy.

3.1.1. Analysis of the white matter adjacent to the deep gray matter

Medium Diffusivity (MD) was the most sensitive metric of white matter difference in HD. MD was increased in the white matter adjacent to the posterior portion of thalamus in the medium-CAP group. The increase was greater and widespread in the high-CAP group (Fig. 4). The MD values for each segment analyzed are reported in the Supplementary Table 2. The source of MD increase was the increase of both radial and axial diffusivity. Fractional anisotropy was not significantly different among groups.

3.1.2. Analysis of the white matter pathways between striatum and cortex

None of the pathways traced by DP between the striatum and cortical areas showed significant difference in DTI between controls and low-CAP or controls and medium-CAP group. Pathways between striatum and many cortical areas had significant MD increase in high-CAP group compared to controls, as shown in Table 2.

3.1.3. Regional analysis of the whole white matter

Once again, MD was the most sensitive DTI metric, showing differences between controls and high-CAP group as shown in Fig. 5. The effect size detected (maximum 8% of MD increase) was smaller than that detected in the analysis of the neighborhood of the deep gray matter (Fig. 4). The most affected areas were the posterior thalamic radiations, corpus callosum, and the peripheral occipital white matter. The MD increase was again due to increases in both radial and axial diffusivities. Although the comparison of MD between medium-CAP group and controls did not reveal significant differences, some areas showed increased MD close to the significance threshold, such as the posterior thalamic radiations. The MD values for each segment analyzed are reported in the Supplementary Table 3. The splenium of the corpus callosum had the lowest FA in the high-CAP group, although no area achieved the significance threshold for FA differences between groups.

3.2. Relationship between shape, DTI metrics, and CAP scores

Fig. 6 shows the trends of shape and MD (y axis) abnormalities according to the CAP scores (x axis), fitted by monotonic curves. There is a positive correlation between the CAP scores and the MD, i.e., MD increases with higher CAP scores. There is a negative correlation between the CAP scores and the amount as each vertex dislocates to match the template (log-Jacobian), i.e., the atrophy increases with higher CAP scores.

4. Discussion

In this study, we combined deep gray matter shape diffeomorphometry and white matter DTI analysis as means of better defining the topography of deep gray and white matter abnormalities in the premanifest stages of HD. We used images from the PREDICT-HD study, which as noted above successfully includes sources of brain MRIs from multiple centers. Across-scanner variations might interfere with the detection of disease-specific structural abnormalities, thereby potentially limiting the use of group analysis collected at several centers. On the other hand, multicenter studies increased generalizability of results, and improved efficiency, particularly for rare or hard-to-recruit cases, such as HD (Paulsen et al., 2008; Tabrizi et al., 2009). Both the algorithm for segmenting subcortical structures and the scalar...
The MD in the white matter surrounding the deep gray matter was observed in the neighborhood of the posterior thalamus in consistently increased close to the disease onset. In addition, higher other groups and hippocampus) and degree in the high-CAP group compared to the pallidus. The atrophy is greater in both location (involving thalamus sively affected over the surface of caudate, putamen, and globus

sectional and longitudinal atrophy in the pre-manifest phase (Aylward et al., 1997, 2000; Rosas et al., 2001; Peinemann et al., 2005; Henley et al., 2009; Vandenbergh et al., 2009; Ginestroni et al., 2010; Georgiou-Karistianis et al., 2013; Sanchez-Castaneda et al., 2013). Other regions, such as the globus pallidus, thalamus and hippocampus, also undergo atrophy, though in smaller magnitude (Paulsen et al., 2010; Aylward et al., 2011; Younes et al., 2014a, 2014b) and mostly after motor the onset. In addition, the measurement of shape has enabled the delineation of regional abnormalities in the striatum and other subcortical structures in the premanifest period (Younes et al., 2014a, 2014b). Reduction of white matter volume was also reported (Tabrizi et al., 2009) in both premanifest and early HD cohorts (Crawford et al., 2013). DTI studies differ regarding specific areas of white matter abnormality in the premanifest stage (Reading et al., 2005; Rosas et al., 2006; Kloppel et al., 2008; Stoffers et al., 2010; Poudel et al., 2014).

In the present study, we integrated the characterization of deep gray matter shape with regional white matter DTI metrics in subjects with premanifest HD to improve the regional mapping of the abnormalities that precede HD onset. Regional abnormalities in the putamen shape, which can be anatomically interpreted as atrophy, occur in the very early stage, with differences being detected between the controls and the low-CAP group. In the medium-CAP group, shape is already extensively affected over the surface of caudate, putamen, and globus pallidus. The atrophy is greater in both location (involving thalamus and hippocampus) and degree in the high-CAP group compared to the other groups.

The MD in the white matter surrounding the deep gray matter was consistently increased close to the disease onset. In addition, higher MD was observed in the neighborhood of the posterior thalamus in the medium-CAP group. According the regional whole brain analysis of the white matter, MD was extensively larger in high-CAP compared to controls, particularly in posterior areas. A trend of increasing MD was seen even in the low-CAP group; in the medium-CAP, areas such as the splenium of the corpus callosum, posterior thalamic radiations, and occipital white matter had MD increase close to the threshold significance. Note that our method was conservative regarding the p-value cutoff, after the permutation and multiple comparisons corrections. This may explain the disagreement with other publications that showed white matter abnormalities occurring at least as early as subcortical gray matter abnormalities (Aylward et al., 2011; Tabrizi et al., 2013).

The tractography analysis revealed increased MD in the topography of streamlines traced between the striatum and multiple cortical areas in the high-CAP group, compared to controls, suggesting that pathology may follow paths of anatomic connections. This needs to be interpreted carefully, since DTI tractography may identify streamlines that imperfectly correspond to actual anatomic connections. Recently, Novak et al. (Novak et al., 2015) showed widespread differences in basal ganglia-cortical structural connectivity in early manifest HD subjects. Variations of DTI indices such as MD and FA may be interpreted as axonal disorganization or axonal degeneration, or defective myelination, or other effects. Our studies cannot define the cellular bases of the changes, but can help map their topography.

The sensitivity of MD or FA to detect group differences depends both on the regional microstructural architecture in question and on the disturbance to this architecture (Oh et al., 2009). In this study, MD was more sensitive than FA in detecting group differences, indicating that HD causes disturbances on the microstructural organization that are more sensitively detected by MD than by FA. Nevertheless, FA tends to be more sensitive to image noise (Pierpaoli and Basser, 1996) as demonstrated in our previous study about MultiCenter Reliability of Diffusion Tensor Imaging (Magnotta et al., 2012), in which among all the DTI scalar measures, FA was the one that had the largest coefficient of variance among all the protocols, although still quite small.
normalities in the white matter, and with a centripetal and posterior tectible atrophy in the deep gray matter would precede detectible ab-

-Section II.5. However, DTI is inherently noisier than T1-WIs, which may result on decreased power on detecting DTI abnormalities compared to

The numbers indicate the MD ratio of high-CAP/controls. Red cells are significant differences between groups, empty cells are pathways that couldn’t be traced, i.e., DP resulted in zero streamlines. The subregions of striatum are ordered according to their anatomical position, from anterior to posterior, as much as possible.

These results, while cross-sectional, rather than longitudinal, may be interpreted in relationship to CAP score (which is a genetic measure of the degree of exposure to the effects of the CAG repeat expansion in HTT), as the cross-sectional reflection of progressive changes in HD. Information from cross-sectional studies such as this one may be useful for guiding therapeutic trials, since subjects in clinical trials will be ascertained based on cross-sectional information and may be selected or stratified based on CAP score or comparable indices. Atrophy started in putamen and caudate, followed by globus pallidum, thalamus, and hippocampus. Our results are consistent with the hypothesis that detectible atrophy in the deep gray matter would precede detectible abnormalities in the white matter, and with a centripetal and posterior gradient of abnormalities in the white matter. There are, however, limitations on comparing white matter microstructural abnormalities and atrophy by imaging. The findings reported in this study may reflect a true biological phenomenon, or just the ability of MRI and of the image analysis employed to detect these abnormalities, or both. Differences in diffeomorphometry and diffusivity between groups were analyzed by the same statistical procedure, via linear effects corrected for multiple comparisons using permutation testing, as detailed in Section II.5. However, DTI is inherently noisier than T1-WIs, which may result on decreased power on detecting DTI abnormalities compared to atrophy. In this sense, our analysis may ensue very conservative results, particularly for DTI. Differences found at conservative group-level

![Fig. 5. Atlas-based white matter DTI analysis. The colors code the increase of MD in areas of significant differences between controls and high-CAP group.](image-url)
analysis, as in this study, can be interpreted with more confidence, and have greater chance to become features on predictive models, which is one of the PREDICT-HD goals, than those found with a less conservative approach. The latter would better indicate small trends at group level and, indirectly, uncover underlying disease mechanisms, which are actually unlikely to be revealed by any MRI study in humans.

Our data suggest that atrophy is detected earliest in the deep gray matter, but that subjacent and closely connected white matter also becomes affected prior to predicted onset. A major current strategy for therapeutics involves lowering levels of mutant htt, via RNA- or DNA-based strategies. One implementation currently in clinical trials involves injection of antisense oligonucleotides into the lumbar CSF and, thus, after CSF circulation, over the cortex. A concern based on our results is that these ASOs may not penetrate to deep gray and deep subcortical structures we demonstrate to be affected during the pre-manifest period. A second potential implementation for clinical trials involves injection of viral vectors into the striatum. Our data suggest that this should be done early (as defined by CAP score), prior to the development of pathology in the deep white matter (thus perhaps even preventing the spread of pathology to the white matter). In addition, it may be necessary to inject the therapeutic agents into the white matter itself.

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.nicl.2016.02.014.

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