Computation of an MRI brain atlas from a population of Parkinson's disease patients

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Abstract. Parkinson’s Disease (PD) is a degenerative disorder of the brain. This study presents an MRI-based brain atlas of PD to characterize associated alterations for diagnostic and interventional purposes. The atlas standardizes primarily the implicated subcortical regions such as the globus pallidus (GP), substantia nigra (SN), subthalamic nucleus (STN), caudate nucleus (CN), thalamus (TH), putamen (PUT), and red nucleus (RN). The data were 3.0 T MRI brain images from 16 PD patients and 10 matched controls. The images used were T1-weighted (T₁₁), T2-weighted (T₂₂), and Susceptibility Weighted Images (SWI). The T₁₁ images were the reference for the inter-subject non-rigid registration available from 3DSlicer. Anatomic labeling was achieved with BrainSuite and regions were refined with the level sets segmentation of ITK-Snap. The subcortical centers were analyzed for their volume and signal intensity. Comparison with an age-matched control group unravels a significant PD-related T₁₁ signal loss in the striatum (CN and PUT) centers, but approximately a constant volume. The results in this study improve MRI based PD localization and can lead to the development of novel biomarkers.

Keywords: Brain 3 Tesla MRI atlas, SWI, Parkinson’s disease biomarkers, striatum, basal ganglia, 3DSlicer, neurodegeneration.

1. Introduction

Parkinson’s disease (PD) is a neurodegenerative disorder that mainly affects the motor system. Magnetic Resonance Imaging (MRI) plays an increasing role as a non-invasive imaging diagnostic tool for neurodegeneration with structural and morphological information. The recent developments of specific sequences such as the Susceptibility-Weighted Imaging (SWI) improves the imaging of midbrain structures significant for PD [1].

The collection and quantification of MRI-datasets enables the assembly of “human brain atlases” as references for spatial normalization and structural segmentation. This can assist not only interventional navigation, but also diagnosis. Many brain atlases are computed from healthy subjects [2,3]. However, general population atlases are typically not representative of the brain degeneration in PD. Thus, recent initiatives promote the construction of disease-specific brain atlases [4].

The aim of this work has been to develop an MRI-based brain atlas representative of PD-brain alterations for interventional and diagnostic purposes. The main goal of the atlas is to localize the implicated subcortical regions, such as the TH, GP, RN, SN, PUT, STN, and CN. This can improve
the focusing of electrode implantation for Deep Brain Stimulation (DBS). Also, the atlas characterizes these anatomic regions in terms of their volumes and signal intensities. The comparison with age-and-gender matched controls can lead to the development of biomarkers for PD.

2. Materials and Methods

The imaging data were from $N = 16$ PD patients of mean age $60 \pm 9$ years (11 M, 5 F). The images $(I_1(x),\ldots,I_N(x))$, where $x$ denotes space, were of three different contrasts, $T_1w$ MPRAGE, $T_2w$, and SWI. The data was retrieved from the Parkinson’s Database of University Medical center Göttingen for acquisitions between 01/2011 and 01/2016. The imaging was performed with a MAGNETOM Trio TIM MR-system 3.0 T (Siemens, Erlangen, Germany) and a dedicated 32-channel head coil. The patients were in total anesthesia. The image analysis was performed with 3DSlicer [5]. The $T_1w$ images were denoised with a median filter. A brain mask, $M_1$, was delineated manually from the corresponding $T_1w$ image. The registration was first intra-subject to the $T_1w$ image. First, a rigid $T_{R,1}$ with 6 degrees of freedom (doF) and then an affine $T_{R,2}$ of 12 doF. The total intra-subject registration is their concatenation, $(T_{R,1} \ast T_{R,2})$, where $\ast$ is convolution. The next step was bias correction with N4ITK [6]. Then, a representative $T_1w$ image was selected as reference, $I_{REF}$ for inter-subject non-rigid registration $T_{NR}$. The $T_{NR}$ was based on BSplines with $6 \times 6 \times 6 = 216$ nodes. All transformations were also applied to $M_1$ to obtain $M_2(T_{NR} \ast T_{R,2} \ast T_{R,1} \ast x)$. The multiplication with $I_1$ gives $I_1 \times M_2(T_{NR} \ast T_{R,2} \ast T_{R,1} \ast x)$ and isolates the brain region. The brain regions histograms of all images were normalized to have the same white matter peak as that of the reference image, $I_{REF}$. Normalization was necessary to make the intensity ranges of the tissues distributions comparable. This allows considering them together and computing their average image. The $T_1w$ brain atlas $I_{atT1}$ was the average image, 
$$I_{atT1} = \frac{1}{N} \sum_{i=1}^{N-1} I_i(T_{NR} \ast T_{R,2} \ast T_{R,1} \ast x).$$

The atlases of the $T_2w$ images, $I_{atT2}$, and SWI images, $I_{atSWI}$, were computed with the application of the corresponding transformations $(T_{NR} \ast T_{R,2} \ast T_{R,1})$ computed from the $T_1w$ images.

To localize the different brain regions of interest we co-registered the $I_{atT1}$ to the labeled BCI-DNI brain atlas available through BrainSuite [7]. This atlas provided initialization of the anatomic regions of interest that were the TH, GP, PUT, and CN. These regions were refined with the level-sets segmentation method of ITK-SNAP [8]. Other fine midbrain structures, namely, the RN, SN and STN were manually segmented by the first author with ITK-SNAP. These segmentations were then corrected by two of the co-authors that are qualified physicians I.E.P. and M.N.P., with three and twelve years of experience respectively. The involvement of two different reviewers following the practices of two different medical centers minimizes subjectivity and thus increases reproducibility. All segmented regions were analyzed for volume and intensity.

The control data consisting of $N = 10$ age-gender matched patients of mean age $48 \pm 5$ years (4 M, 6 F) without neurodegenerative changes were processed with the same steps as those described above. They produced a $T_1w$ brain atlas $I_{atT1}$ as well as the anatomic regions of the TH, GP, PUT, and CN. The information derived from the segmentations of the brain structures for PD and healthy subjects were compared in terms of their volume and their intensities with Student’s 2-tailed t-test or with the non-parametric Mann–Whitney U tests with Sigma Plot™ (Systat Software Inc., San Jose, CA USA). The diagnostic sensitivity (SE) and specificity (SP) of the analyzed parameters were evaluated with Receiver Operating Characteristics (ROC) and the Cut-off values were determined with Youden J statistics.

3. Results

In this study both signal intensity and volume have been selected as MRI correlates for the pathological changes occurring in PD. Neurodegeneration is intuitively associated with loss of
volume, “involution”, of the subcortical nuclei. Moreover, depositions of protein inclusions, cell debris and gliosis significantly modify the proton density and variously affect the $T_1w/T_2w$ signal [9].

3.1. MRI-based brain atlases, Volume and Signal Intensity of the basal ganglia as biomarkers

The images from N=16 PD patients and N=10 control provided brain atlases for the PD-brain and for age- and gender-matched controls as shown in ‘figure 1’. In contrast to the midbrain tegmentum (SN and RN), the GP, the striatum which is part of basal ganglia, and the TH are well identifiable in the $T_1w$ MPRAGE image from the 16 PD and the 10 control subjects. Based on the segmentations outer contours we generated 3D-surface renderings with 3DSlicer for the striatal area focusing on the CN ‘figure 2’ (i, ii, iii). Representative axial, sagittal and coronal $T_1w$ images with color-coding of the segmented nuclei (blue) are shown in the panel below the surface renderings in ‘figure 2’ (iv, v, vi). PD patients were analyzed for volume and $T_1w$ signal intensity changes compared to the control group.

We analyzed the CN whose volume is not significantly affected by PD, $P=0.43$, 2-tailed t-test in ‘figure 2’ (b), most of the other analyzed regions followed the same pattern. However, the $T_1w$ signal intensity is significantly reduced in PD patients in ‘figure 2’ $P < 0.0001$, 2-tailed t-test (c) by approximately 8% from the average control value. The parameter for the ROC analysis was the $T_1w$ intensity and its diagnostic efficacy is high as shown in ‘figure 2’ (d), with an $AUC = 0.93 \pm 0.07$. By implementing Youden statistics for the $T_1w$ signal intensity we could reveal a cut-off diagnostic value of 352.5 with a SE of 100% and a SP of 90% for PD versus the control shown in ‘figure 2’ (c). In summary, even though PD is not associated with striatal involution, it affects the $T_1w$ signal intensity of the caudate nucleus compared to an age- and gender- matched control population. Thus, the ROC analysis suggests that PD-related signal loss in the $T_1w$ sequence is a potential sensitive diagnostic parameter for PD.

![Figure 1](image1.png)  
**Figure 1 (left).** MRI brain atlas. (I) Probabilistic average images from population, $T_1w$, N=10. (II) Probabilistic average images from PD patients, $T_1w$. (III) $T_2w$ PD and (IV) SWI PD, N=16.  
**Figure 2 (right).** Volume and intensity analysis of the caudate nucleus. Sample PD-brain 3D surface-renderings of the basal ganglia (CN, dark green; PUT, light pink; GP, light green; TH, dark pink) and the midbrain nuclei (SN, marine blue; RN, red) as well as the STN (black). (Av) Axial, (Av) sagittal and (Avi) coronal $T_1w$ sections with blue-shaded CN. (B) Volume analysis of the CN (C) Intensity analysis (D) ROC of the $T_1w$ MPRAGE intensity of the CN, PD N=16, control N=10.
4. Discussion

The PD brain atlas and quantification in this study focuses not only on the primarily affected midbrain regions (SN), but also on the downstream affected basal ganglia, especially the striatum. The striatum, receives dopaminergic innervation from the SN, hence undergoing “denervation” secondarily to nigral neurodegeneration [10]. This is associated with neural dystrophy but not necessarily with neuronal death [11]. This is in agreement with our findings, classical radiological descriptions and neuropathology converge toward the opinion that PD is not associated with striatal involution [12]. Apart from the loss of volume, we analyzed the subregions for signal intensity as an imaging correlate of histological remodeling. We showed a significant signal loss of the striatum (CN) in PD compared to controls. Such signal alteration may represent initial structural changes and neuronal loss that do not necessarily reflect volume loss.

The main target of this study was to create an MRI brain atlas of the PD brain based on retrospective data acquisition from an existing database. The platform of the atlas is flexible to modifications and can be enriched with additional subjects. A limitation of this study is that the control database consisted of only $T_1w$ images. Thus, volume and intensity alterations could not be reported for structures defined by the $T_2w$ image and the SWI signal, specifically for the SN and STN.

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