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Nigrosome-1 on Susceptibility Weighted Imaging to Differentiate Parkinson’s Disease From Atypical Parkinsonism: An In Vivo and Ex Vivo Pilot Study

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Summary

Background:
Previous case-control studies have suggested that the absence of a swallow-tail appearance in the substantia nigra on high-resolution SWI, representing nigrosome-1, has high accuracy to identify Parkinson’s disease (PD). The first goal of our study was to evaluate nigrosome-1 ex vivo using optimized high-resolution susceptibility sensitive MRI. Our second goal was to evaluate its diagnostic value in vivo using a clinical 3T SWI sequence to differentiate between PD and atypical parkinsonism (AP) in a cohort of patients with early-stage parkinsonism.

Material/Methods:
Case-control pilot study to evaluate nigrosome-1 ex vivo (2 PD, 2 controls), using high-resolution susceptibility sensitive sequences at 11.7 T MRI. Next, evaluation of nigrosome-1 in vivo using a clinical 3 T SWI sequence in a prospective cohort of 60 patients with early-stage parkinsonism (39 PD, 21 AP). Moreover, 12 control subjects were scanned.

The bilateral substantia nigra was evaluated by two neuroradiologists for the presence, absence or indecisive presence of nigrosome-1. The discriminatory power was evaluated by Receiver-Operating Characteristic.

Results:
We identified nigrosome-1 in ex vivo control subjects. Nigrosome-1 was not identified in the ex vivo PD cases. In our prospective clinical cohort study, the AUC for the swallow-tail sign to discriminate between PD and AP was 0.56 (0.41–0.71) for reader 1 and 0.68 (0.55–0.82) for reader 2.

Conclusions:
The diagnostic accuracy of the swallow-tail sign was marginal to discriminate between PD and AP using our clinical 3 T SWI sequence.

MeSH Keywords: Brain Diseases • Magnetic Resonance Imaging • Parkinson Disease • Parkinsonian Disorders

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Background

Parkinson’s disease (PD) is a neurodegenerative disorder with incidence increasing with age. The diagnosis of PD is made on clinical grounds, based on classic motor symptoms including bradykinesia, rest tremor or rigidity [1,2]. Conventional brain MRI lacks a diagnostic marker specific for PD [3]. Previously, studies have been published with promising results regarding T1 and T2 signal intensity changes of the substantia nigra (SN) in PD [4,5], which could not be reproduced in follow-up studies. In the diagnostic work-up of parkinsonism, brain MRI is therefore mainly used to exclude other, more rare causes of parkinsonism such as vascular damage or hydrocephalus. Also, abnormalities on brain MRI could support the diagnosis of a neurodegenerative disorder other than PD, usually referred to as atypical parkinsonism (AP). These include multiple system atrophy (MSA), progressive supranuclear palsy (PSP), dementia with Lewy bodies (DLB) and corticobasal syndrome (CBS).

New MRI techniques such as diffusion tensor imaging (DTI) and magnetization transfer imaging (MTI) may have the potential to provide a new diagnostic marker specific for PD [6], but clinical application of these techniques is limited because study results are not consistent and validated diagnostic criteria are generally lacking [6,7]. This also accounts for quantitative measures of iron content of the SN in PD, demonstrated as increased susceptibility on T2* or Susceptibility Weighted Imaging (SWI) sequences [8–10].

In recent in vivo and postmortem studies using high-resolution susceptibility sensitive MR imaging techniques (7 T but also 3 T MRI), a new diagnostic marker has been reported with high accuracy to identify PD [11–13]. A subregion in the ventrolateral part of the substantia nigra lacking susceptibility and bordered by two lines of susceptibility, resembling a swallow-tail (Figure 1), represents nigrosome-1 and is identified in healthy controls. The absence of this swallow-tail configuration has been reported in PD, with observed good discrimination between PD and control subjects (reported accuracy 91–96%) [14]. It is not known whether the absence of the swallow-tail sign on SWI could discriminate PD from AP.

The first goal of our pilot study was to evaluate the swallow-tail appearance in the substantia nigra ex vivo, using optimized high-resolution susceptibility sensitive MRI sequences. Our second goal was to evaluate the diagnostic value of the swallow-tail sign to differentiate between PD and AP in a cohort of patients with early-stage parkinsonism using a clinical 3 T SWI sequence.

Material and Methods

Ex vivo case-control study (11.7 T MRI)

Formalin-fixed postmortem midbrain samples of 2 patients with histopathologically confirmed PD (76 y/o male and 81 y/o female) and 2 controls (76 and 84 y/o females without a neurodegenerative disorder) were collected. Slices of the midbrain including the lower part of the substantia nigra were rehydrated with phosphate buffered saline for one week to reduce fixation-induced T2 shortening. A unilateral part of the midbrain samples (measuring approximately 10×10×5 mm), was placed in a syringe filled with Galden D40 perfluoropolyether (Solvay Solexis, New York). The samples were stored at room temperature for 72 hours prior to scanning and air bubbles were removed. This method is in accordance with a previous ex vivo MRI study of our group [15].

Imaging was performed on an 11.7-T BioSpec Avance III small animal MRI system (Bruker BioSpin, Ettlingen, Germany) equipped with an actively shielded gradient set of 600 mT/m and operated by the ParaVision 5.1 software platform. A circular polarized volume resonator was used for signal transmission, and an actively decoupled mouse brain quadrature surface coil with integrated combiner and preamplifier (Bruker BioSpin) for signal reception. After standard adjustments, 2D and 3D gradient echo sequences were acquired. Details of the scanning protocol are provided in Table 1.

The images were visually inspected by two neuroradiologists (FJAM and BG) for the presence of a swallow-tail appearance in the substantia nigra pars compacta, in order to identify nigrosome-1.

Clinical cohort and case-control study (clinical 3T SWI)

A prospective observational study of 60 patients presenting with early stage parkinsonism, part of a larger clinical cohort study, was performed. Those patients were prospectively recruited as part of a larger clinical cohort study. The work described was carried out in accordance with The Code of Ethics of the World Medical Association (Declaration of Helsinki) for experiments involving humans. The medical ethics committee of our hospital
Approved the study and all participants gave written informed consent prior to inclusion. Patients were recruited at our outpatient movement disorder clinic in the period 2010–2012.

Study inclusion criteria were clinical signs and symptoms of parkinsonism (hypokinetic-rigid syndrome) with uncertain clinical diagnosis and disease duration of less than 3 years. Exclusion criteria were age under 18 years, prior brain surgery, presence of another neurodegenerative disorder and unstable co-morbidity.

After a mean clinical follow-up of 24.6 (±12.4) months, 'silver standard' diagnoses could be made by two experienced clinicians (AR, RE): 39 patients were diagnosed with PD and 21 patients with AP (13 MSA-P, 3 PSP, 3 DLB, 1 vascular parkinsonism, 1 CBS). Those diagnoses were made according to international diagnostic criteria [16–21], based on neurological signs that developed during the course of the disease (as identified during repeat neurological exams), rate of disease progression and treatment response. In addition, 12 controls were included. Demographic criteria of the study groups are summarized in Table 2. In comparison to PD, disease severity and severity of motor symptoms were higher for AP (p<0.05). After a mean clinical follow-up of 24.6 (±12.4) months, 'silver standard' diagnoses could be made by two experienced clinicians (AR, RE): 39 patients were diagnosed with PD and 21 patients with AP (13 MSA-P, 3 PSP, 3 DLB, 1 vascular parkinsonism, 1 CBS). Those diagnoses were made according to international diagnostic criteria [16–21], based on neurological signs that developed during the course of the disease (as identified during repeat neurological exams), rate of disease progression and treatment response. In addition, 12 controls were included. Demographic criteria of the study groups are summarized in Table 2. In comparison to PD, disease severity and severity of motor symptoms were higher for AP (p<0.05).

At baseline of this study, all patients had a 3-T brain MRI scan (Magnetom Trio, Siemens, Erlangen, Germany), which included a clinical SWI sequence with a high in-plane resolution but 3-mm slice thickness (0.63×0.63×3 mm voxels). Details of the scanning protocol are provided in Table 1. These SWI sequences were visually evaluated by two neuroradiologists (FJAM and SS), blinded to clinical information and diagnoses. The bilateral substantia nigra was scored for the unambiguous presence, unambiguous absence, or indecisive presence of the swallow-tail sign. Unilateral absence of the swallow-tail sign was considered indicative of PD irrespective of the other side, as the onset of PD can be asymmetrical.

Cohen’s kappa co-efficient was used to evaluate intra- and inter-rater variability. Agreement was graded as: kappa <0.20, poor agreement; 0.21–0.4, fair agreement; 0.41–0.60, moderate agreement; 0.61–0.80, good agreement; >0.80, excellent agreement. One reader (FJAM) scored all MRI studies twice with a two-week interval in order to evaluate the intra-rater variability.

The discriminative power of the swallow-tail appearance was evaluated for both readers with the Receiver-Operating-Characteristic (ROC).

### Results

**Ex vivo case-control study (11.7 T MRI)**

Nigrosome-1 was identified by a swallow-tail appearance in the ventrolateral substantia nigra pars compacta on the MR images of the two control samples, while nigrosome-1 was not identified in the two PD samples (Figure 2).

<table>
<thead>
<tr>
<th>Study/MR field strength</th>
<th>Sequence</th>
<th>TR/TE</th>
<th>Flip angle*</th>
<th>Matrix size</th>
<th>Voxel (mm)</th>
<th>No. of averages</th>
<th>Acquisition time</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ex vivo/11.7 T</strong></td>
<td>2D MGE*</td>
<td>1248</td>
<td>12</td>
<td>512×512</td>
<td>0.079×0.079×0.64</td>
<td>2</td>
<td>16 min</td>
</tr>
<tr>
<td></td>
<td>3D MGE* with 11 echos</td>
<td>57/3.4–53.4</td>
<td>30</td>
<td>256×256</td>
<td>0.112×0.112×0.112</td>
<td>–</td>
<td>47 min</td>
</tr>
<tr>
<td><strong>In vivo SWI 3 mm sliced/3 T</strong></td>
<td>3D gradient echo SWI</td>
<td>29/20</td>
<td>15</td>
<td>384×384</td>
<td>0.63×0.63×x3</td>
<td>1</td>
<td>4 min, 42 sec</td>
</tr>
</tbody>
</table>

* MGE – multi gradient echo.

<table>
<thead>
<tr>
<th>Table 1. Details of the MRI scanning protocols used.</th>
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<tr>
<td>Study/MR field strength</td>
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<tr>
<td><strong>Ex vivo/11.7 T</strong></td>
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* MGE – multi gradient echo.

<table>
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<tr>
<th>Table 2. Demographic data of the prospective study group.</th>
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<tbody>
<tr>
<td>Study/Group</td>
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<tr>
<td>-------------</td>
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<tr>
<td>Mean age (yrs)</td>
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<tr>
<td>Gender (M: F)</td>
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<tr>
<td>Disease duration (months)</td>
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<tr>
<td>MMSE</td>
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<td>H&amp;Y</td>
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<td>UPDRS-III</td>
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</tbody>
</table>

PD – Parkinson’s disease; AP – atypical parkinsonism; MMSE – mini-mental state examination; H&Y – Hoehn and Yahr staging scale; UPDRS-III – Unified Parkinson Disease Rating Scale – III. * Student’s t-test p-value <0.05.
Clinical cohort and case-control study (clinical 3 T SWI)

Table 3 demonstrates the distribution of nigrosome-1 as identified by both readers in PD, AP and controls. Inter-rater variability proved to be fair (kappa 0.35). Intra-rater variability proved to be moderate (kappa 0.44). For reader 2, nigrosome-1 was identified in the majority of PD patients (56%), while this was the case in a minority of PD patients for reader 1 (23%).

The ability of the swallow-tail sign to discriminate between PD and controls resulted in an AUC of 0.72 (0.58–0.87) for reader 1 and 0.58 (0.39–0.77) for reader 2. For the discrimination between PD and AP, the AUC was 0.56 (0.41–0.71) for reader 1 and 0.68 (0.55–0.82) for reader 2.

Table 3. The absence, presence and indecisive presence of nigrosome-1 in Parkinson's disease, atypical parkinsonism and controls.

<table>
<thead>
<tr>
<th></th>
<th>Nigrosome-1</th>
<th>Nigrosome-1</th>
<th>Nigrosome-1</th>
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<tbody>
<tr>
<td></td>
<td>absent</td>
<td>present</td>
<td>indecisive</td>
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<tr>
<td>Reader 1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PD (n=39)</td>
<td>23 (59%)</td>
<td>9 (23%)</td>
<td>7 (18%)</td>
</tr>
<tr>
<td>AP (n=21)</td>
<td>14 (67%)</td>
<td>6 (29%)</td>
<td>1 (5%)</td>
</tr>
<tr>
<td>Controls (n=12)</td>
<td>1 (8%)</td>
<td>8 (67%)</td>
<td>3 (25%)</td>
</tr>
<tr>
<td>Reader 2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PD (n=39)</td>
<td>13 (33%)</td>
<td>22 (56%)</td>
<td>4 (10%)</td>
</tr>
<tr>
<td>AP (n=21)</td>
<td>14 (67%)</td>
<td>7 (33%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Controls (n=12)</td>
<td>6 (50%)</td>
<td>5 (42%)</td>
<td>1 (8%)</td>
</tr>
</tbody>
</table>

PD – Parkinson’s disease; AP – atypical parkinsonism.

Clinical cohort and case-control study (clinical 3 T SWI)

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Discussion

A compartmental organization of the substantia nigra, consisting of the nigral matrix and nigrosomes, was first described based on the immunohistochemical staining of calbindin in striatonigral afferent fibers [22]. It was reported that the largest of the nigrosomes called nigrosome-1, located in the ventro-lateral substantia nigra, was most affected in PD exhibiting the maximum depletion of dopaminergic cells [23]. Based on the results of previous case-control studies [11–14], the swallow-tail sign seems to be a promising new diagnostic MRI marker in the diagnostic work-up of parkinsonism. In contrast to quantitative MRI techniques, this possible new diagnostic marker can be easily implemented in everyday clinical practice because there is no requirement for complicated post-processing prior to visual assessment. Using an 11.7-T MRI scanner, we were able to identify the swallow-tail appearance of nigrosome-1 in the substantia nigra pars compacta in ex vivo midbrain samples of controls. In two midbrain samples with histopathologically confirmed PD, this structure was absent. Yet, no studies have been published evaluating nigrosome-1 in AP, and evaluating its diagnostic value in discriminating PD from AP.

In our prospective observational cohort study, the diagnostic accuracy of the swallow-tail sign to differentiate between PD and AP was marginal, with limited intra- and inter-rater agreement. This also accounts for the differentiation between PD and controls, with a diagnostic performance clearly lower than the case-control studies published previously. A likely explanation for this discrepancy is that the 3-mm slices of our clinical SWI protocol were too thick to reliably visualize nigrosome-1, despite its high in-plane resolution. This explanation is supported by the fact that a swallow-tail configuration in the substantia nigra was missing in a significant part of the control subjects on this sequence. The minimal required spatial resolution of SWI to reliably identify nigrosome-1 has not been reported, though an optimized 3D high-resolution SWI is probably crucial with increased confidence at higher magnetic field strengths. In a very recent case-control study comparing 7 T and 3 T for the evaluation of nigrosome-1 [23], the confidence in revealing the inner structure of the substantia nigra on 3 T was inferior to that of 7 T as demonstrated by lower intra- and inter-observer agreement at 3 T. Also, the diagnostic accuracy to identify PD proved to be slightly lower for 3 T (86%) in comparison to 7 T MRI (96%). Increased T2* contrast and magnetic susceptibility effects of paramagnetic substances at higher magnetic fields is the explanation the authors consider the most likely for the superior imaging performance of 7T, rather than a higher spatial resolution which only differed slightly between 3 T and 7 T [24].

Furthermore, it can be debated whether a swallow-tail configuration identified on our relatively thick-sliced SWI sequence actually represents nigrosome-1 in the inner structure of the substantia nigra, or would be a reflection of the closely related substantia nigra and subthalamic nucleus. This bears the risk of a false-positively identified nigrosome-1.

Another discrepancy with previous studies is that a swallow-tail configuration in the substantia nigra was frequently seen in our prospective cohort of patients diagnosed with PD. Our PD patients were probably scanned earlier in the course of the disease (mean disease duration of 21.6 months) than the case-control studies published previously (which included patients with disease duration of up to 10 years) [12,14,24]. The absence of nigrosome-1 in the substantia nigra in PD on SWI is probably not explained by predominant degeneration or volume loss, but is most likely explained by pathologically increased iron accumulation with increased susceptibility on SWI as a result [14]. However, in previous studies increase of the iron concentrations of the substantia nigra did not seem to
be related to the disease duration in PD, while it was correlated with the severity of motor symptoms [25,26]. Another possible explanation for the absence of nigrosome-1 in PD is decreased neuromelanin content of the substantia nigra with decreased iron storage capacity leading to more free iron with paramagnetic properties [27]. It needs to be determined whether the absence of nigrosome-1 on SWI in PD is related to disease duration and whether it correlates with loss of presynaptic dopamine transporters, as demonstrated by nuclear scan techniques. In case the absence of nigrosome-1 on SWI was not related to disease duration, it could prove to be a useful diagnostic MRI marker in the work-up of early-stage parkinsonism. It needs to be determined whether it is PD-specific and not valid for atypical parkinsonism, as absence of the swallow-tail sign was also noted in AP in our clinical cohort (Figure 3).

There are some limitations to our study. Our in vivo and ex vivo case-control studies included only a few subjects. Because case-control studies with a larger amount of subjects have been published recently, we only aimed to reproduce the identification of the healthy nigrosome-1, and its absence in PD. A limitation of our clinical prospective observational cohort is that we do not have any post-mortem confirmation of the diagnoses, and therefore we cannot fully rule out misdiagnosis. Well-designed prospective clinical cohort studies are warranted for assessing the diagnostic accuracy of the swallow-tail sign in early-stage parkinsonism. Conducting such a study is challenging because obtaining histopathological confirmation for larger study populations is practically impossible. Although high rates of misdiagnosis have been reported on for the clinical diagnosis, pathological studies show high accuracy levels (>90%) for the clinical diagnosis when the diagnosis was made by a movement disorder specialist after a minimal follow-up of 2 years [28].

Conclusions

The diagnostic accuracy of the swallow-tail sign was marginal to discriminate between PD and AP using our clinical 3 T SWI sequence with high in-plane resolution but 3-mm slice thickness. The diagnostic value of the swallow-tail sign in the work-up of early-stage parkinsonism using an optimized high-resolution SWI sequence needs to be determined in future well-designed prospective clinical cohort studies.

Acknowledgements

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References:


