Visualization of Nigrosome 1 from the Viewpoint of Anatomic Structure


*AJNR Am J Neuroradiol* 2020, 41 (1) 86-91
doi: [https://doi.org/10.3174/ajnr.A6338](https://doi.org/10.3174/ajnr.A6338)
[http://www.ajnr.org/content/41/1/86](http://www.ajnr.org/content/41/1/86)
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ABSTRACT

BACKGROUND AND PURPOSE: Parkinson disease is related to neurodegeneration and iron deposition in the substantia nigra pars compacta and nigrosome 1. However, visualization of nigrosome 1 via MR imaging is poor owing to the bilateral asymmetry, regardless of whether it is healthy. We focused on the magic angle and susceptibility effect and evaluated the anatomic slant structure of nigrosome 1 by tilting subjects’ heads in the B0 direction.

MATERIALS AND METHODS: To investigate the effectiveness of the magic angle, we tilted the volunteers’ heads to the right and left in the B0 direction or not at all for evaluating correlations between the degree of head tilting and visualization of the right nigrosome 1 and left nigrosome 1 using 3D spoiled gradient-echo sequences with multiecho acquisitions. We evaluated the susceptibility of nigrosome 1 and the local field using quantitative susceptibility mapping to assess static magnetic field inhomogeneity.

RESULTS: The heads tilted to the right and left showed significantly higher contrasts of nigrosome 1 and the substantia nigra pars compacta than the nontilted heads. No significant differences were observed in the visualization and susceptibility between the right nigrosome 1 and left nigrosome 1 for each head tilt. The effect of the magic angle was remarkable in the nontilted heads. This finding was supported by quantitative susceptibility mapping because the anatomic slant structure of nigrosome 1 was coherent between the axis of nigrosome 1 and the magic angle.

CONCLUSIONS: The asymmetric visualization of nigrosome 1 is affected by the magic angle and susceptibility. The anatomic slant structure of nigrosome 1 causes these challenges in visualization.

ABBREVIATIONS: PD = Parkinson disease; SNc = substantia nigra pars compacta

Parkinson disease (PD) is associated with major pathologic degeneration of dopaminergic neurons, mainly in the basal ganglia and especially in the substantia nigra pars compacta (SNc). It has been reported that PD or other neurodegenerative disorders include iron accumulation in the SNc. In addition, normally, the iron load in the basal ganglia increases with age.

Received June 29, 2019; accepted after revision October 10.
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The sponsor of this research is Hitachi Ltd.
Paper previously presented at: Annual Meeting of the European Congress of Radiology, March 1–5, 2017; Vienna, Austria (C-1756); and Annual Meeting of the Japanese Society of Radiological Technology, April 13—16, 2017; Yokohama, Japan.
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http://dx.doi.org/10.3174/ajnr.A6338
signals of nigrosome 1 commonly diminish; this feature reflects iron accumulation and neurodegeneration. However, the MR imaging visualization of nigrosome 1 is often poor because of the asymmetry of this cell cluster, regardless of whether nigrosome 1 is healthy. In addition, little attention has been paid to poor visualization due to asymmetry. This is important because poor visualization could result in misclassification of a healthy or abnormal nigrosome 1.

In this study, we explored the cause of nigrosome 1 being poorly visualized despite healthy conditions. We focused on the magic angle and the susceptibility effect, and we evaluated the anatomic slant structure (head tilt) of nigrosome 1 in the B0 direction using combined multi-gradient-echo images and by tilting the subjects’ heads in the B0 direction.

**MATERIALS AND METHODS**

**Subjects**

Fourteen young healthy volunteers (3 women and 11 men; mean age, 25 years; ranging from 22 to 31 years) participated in the study. This study was approved by the institutional review board at Nagoya City University. The volunteers participated in the study of their own free will, and informed consent was obtained from all the participants.

**Data Analysis**

The volunteers’ heads were tilted to the right (20.9° ± 5.8°) or left (18.9° ± 6.6°) in the B0 direction or not at all, to investigate the effectiveness of the magic angle (Fig 1). The 3D spoiled gradient-echo sequences with multiecho acquisitions targeted the midbrain and were oriented orthogonally across from the anterior midbrain (Fig 2A, -B). We evaluated the correlations between the degree of head tilting and visualization of the right nigrosome 1 and left nigrosome 1. To calculate the contrast of nigrosome 1 and the remaining SNc that was positioned around nigrosome 1 to assess its visualization, we used the following equation:

\[
\text{Contrast} = \frac{S_{I1} - S_{SNc}}{S_{I1} + S_{SNc}},
\]

where \(S_{I1}\) and \(S_{SNc}\) are the signal intensities of nigrosome 1 and SNc, respectively (Fig 2C). We calculated the SDs of the contrasts of the right nigrosome 1 and left nigrosome 1 and the SNc for each head tilt, to evaluate the reproducibility of contrast measurement. The contrast was measured 6 times in 1 volunteer using right head tilting and left head tilting. We used ImageJ 1.48 image-processing software (National Institutes of Health, Bethesda, Maryland) to analyze the images.

In addition, we evaluated the susceptibility of nigrosome 1 and the local field using quantitative susceptibility mapping\(^{16,17}\) to assess the visualization of nigrosome 1 changed by the phase modulation due to the dipole effect\(^{18-20}\). To estimate the tissue-generated local field map and the susceptibility map, we used the multi-spoiled gradient-echo sequence in healthy volunteers. First, the local field map was estimated from the acquired multiphase images by means of sophisticated harmonic artifact reduction using the phase data method with variable kernel sizes from 0.75 at the boundary of the brain to 30 mm toward the

| Table 1: Number of visualizations of discriminable nigrosome 1 when the subjects’ heads were tilted in the B0 direction (n = 14) |
|---|---|---|---|---|
| **Tilting Head in B0 Direction** | **Right** | **Left** | **None** | **Right** | **Left** |
| Nigrosome 1 | 9 | 9 | 6 | 5 | 10 | 11 |
| Discriminable nigrosome 1 (No.) | | | | | | |
center of the brain.\textsuperscript{21,22} Then, the susceptibility map was reconstructed with an improved sparse linear equation and the least-squares algorithm.\textsuperscript{23} All data were processed with the in-house Matlab R2018a (MathWorks, Natick, Massachusetts) program.

**MR Imaging Acquisition**

All examinations were performed on a 3T MR imaging system (Trillium Oval; Hitachi, Tokyo, Japan) with a 15- or 32-channel head matrix coil. The evaluation was based on a combination of 3D spoiled gradient-echo images with 9 echoes. To combine the 3D spoiled gradient-echo with each TE magnitude, we used the following parameters: TR, 83 ms; minimum and maximum TE, 18.5 and 55.3 ms, respectively (number of combined echoes, 9; echo spacing, 4.6 ms); flip angle, 10°; section thickness, 1.5 mm; matrix, 300 × 300; FOV, 190 mm (in-plane resolution, 0.63 × 0.63 mm); parallel imaging factor, 1.1; number of signal averages, 1; receiver bandwidth, 210 kHz; and acquisition time, 4 minutes 53 seconds. The receiver bandwidth was fixed for each TE in the multiecho imaging technique. Quantitative susceptibility mapping was performed using the following parameters: TR, 34 ms; TE, 6-30 ms at 6-ms intervals; flip angle, 15°; section thickness, 1.0 mm; matrix, 192 × 192; FOV, 192 mm (in-plane resolution, 1.0 × 1.0 mm); parallel imaging factor, 1.9; number of signal averages, 1; receiver bandwidth, 77 kHz; and acquisition time, 10 minutes 5 seconds.

**Statistical Analysis**

A Wilcoxon signed rank test with a Bonferroni correction for non-normally distributed data was performed to examine the relationship between the visualization of nigrosome 1 and the degree of head tilting to the right and left. Statistical analysis was performed using SPSS (IBM, Armonk, New York). $P < .05$ and $P < .017$ (.05/3 after Bonferroni correction based on 3 comparisons per task) were considered statistically significant.

**RESULTS**

The visualization of nigrosome 1 depends on the difference in the field strength; however, it should be visualized bilaterally,
especially when visualizing a healthy nigrosome 1. Poor visualization due to asymmetry of the bilateral nigrosome 1 was discovered, though all the subjects in this study were young and healthy and had no nucleus degeneration. Additionally, in some subjects, we were unable to find a discriminable nigrosome 1 (Table 1).

The heads tilted to the right and left showed a significantly higher contrast of nigrosome 1 and SNc than the nontilted heads (Fig 3). Figure 4 shows examples of the visualization of nigrosome 1 with a right head tilt (A), without a head tilt (B), and with a left head tilt (C). No significant differences were observed in the visualization and susceptibility between the right nigrosome 1 and left nigrosome 1 for each head tilt. The contrast of nigrosome 1 and SNc showed sufficient reproducibility among the 6 measurements (Table 2). However, the effect of the magic angle was remarkable in the nontilted heads and was also supported by quantitative susceptibility mapping. The boundary of nigrosome 1 and SNc is likely visible (Fig 5A, B, C), because according to the anatomic slant structure of nigrosome 1 along the magic angle, the dipolar interaction has angular dependence (Fig 5C).

**DISCUSSION**

Conventional MR imaging characteristics of the substantia nigra in PD involve increased iron-related contrast enhancement, which reflects the sensitivity of the gradient-echo sequences to the resulting changes in $R_2^*$ relaxivity—that is, the signal intensity on a 3D spoiled gradient-echo is diminished specifically in the lateral portion of the SNc. Iron is the main source of susceptibility in the regions of the nucleus in the midbrain. In addition, a boundary is visible between nigrosome 1 and SNc on gradient-echo images because they naturally have different degrees of iron deposition. The important point here is that the degree of iron deposition is lower than that of neurodegeneration in the midbrain.

In principle, one would expect to visualize a healthy nigrosome 1 bilaterally on MR imaging, but the visualization is often poor because of its asymmetry. Furthermore, SNc is also asymmetric. Hence, a healthy nigrosome 1 and an abnormal nigrosome 1 might be misclassified. The problem that we have assessed here (ie, misclassification of a healthy nigrosome 1 and an abnormal nigrosome 1) has seldom been studied.

The differences in visualization were due to the way in which the subjects’ heads were tilted (ie, to the right or left), which caused asymmetric visualization of nigrosome 1. In our study, we found that when the heads were tilted to the right or left, the contrast between nigrosome 1 and SNc on the images was significantly higher than that with the nontilted heads. Furthermore, the results in Fig 4 strongly suggest the reproducibility of contrast measurements by manually drawing the ROIs and measuring the signal intensities of nigrosome 1 and SNc. We focused on the magic angle in the $B_0$ direction. The magic angle that occurs at approximately 54.7° in the $B_0$ direction and so forth along the axis of the magic angle24,25 accounts for interference with this boundary in the nontilted heads. The relationship between the poor visualization of nigrosome 1 and the magic angle causes asymmetric visualization of nigrosome 1 (Table 1 and Fig 3).

We evaluated the anatomic slant structure of nigrosome 1 using quantitative susceptibility mapping, which enables quantitative investigations of the iron content in tissues.17,21,26–29 $R_2^*$ increases exponentially with increasing age,7,8 but all the subjects included in this study were young and healthy without any nucleus degeneration. Hence, nigrosome 1 and the local field were...
not assessed with susceptibility-weighted imaging and the T2* image technique; instead, quantitative susceptibility mapping, which is more sensitive, was used. The results indicated that the anatomic slant structure of nigrosome 1 was coherent between the axis of nigrosome 1 and the magic angle (Fig 5).

Quantitative susceptibility mapping can also be used for the detection and quantification of iron deposition, whereas the status of the nuclear regions of the midbrain is assessed through diffusion tensor imaging. The bulk tissue microstructure can be measured with this technique on the basis of the tissue microenvironment with mean diffusivity and fractional anisotropy. The mean diffusivity increases with microscopic barrier disruption and extracellular fluid accumulation, and fractional anisotropy provides information on the microstructural integrity of highly oriented microstructures (eg, myelin).30,31 MR diffusion kurtosis imaging has recently been proposed as a means of quantifying non-Gaussian water diffusion; in general, the regions of high fractional anisotropy are also characterized by increased mean kurtosis.

With regard to PD, previous diffusion tensor imaging studies on affected patients have demonstrated a decrease in fractional anisotropy in SNc compared with the fractional anisotropy in SNc obtained in studies conducted on healthy control subjects.32,33 Patients with PD exhibit significantly higher R2* values in the SNc, lower fractional anisotropy values in the SNc and thalamus, and higher mean diffusivity values in the thalamus than control subjects.1 However, the appearance of artifacts, which are more common in diffusion MR imaging sequences, depends on the local magnetic field.34 Inhomogeneities in the local magnetic field, such as those induced by iron deposition, lead to some artifacts in diffusion MR imaging sequences and have an effect on ADC. The sensitivity of the magnetic susceptibility changes because motion-probing gradients into echo-planar imaging are used in diffusion MR imaging sequences. Hence, it is possible that ADC is underestimated in the presence of iron depositions. Conversely, a 3D spoiled gradient-echo with a multi-echo technique is more robust for evaluating nigrosome 1 than a diffusion MR imaging sequence in local magnetic field inhomogeneities. However, as mentioned earlier, the problem with the magic angle in the B0 direction affecting nigrosome 1 visualization became clear in this study.

There were some limitations to this study. The local field gradients by blood containing deoxyhemoglobin were paramagnetic.35 In the future, we will be able to evaluate the magnetic susceptibility of each tissue with more focus. Moreover, the effect of aging needs to be examined in detail. It has been reported that the mean diffusivity and fractional anisotropy are highly influenced by physiologic aging.36 Therefore, we will evaluate older subjects who are added to the study. The current MR imaging technique has made it possible to use appropriate parameters in this study (eg, the number or timing of combined echoes).

**CONCLUSIONS**

The visualization of nigrosome 1 is affected by the magic angle, thus causing asymmetric visualization. We observed that visualization was improved when the head was tilted to the right and left in the B0 direction. Furthermore, nigrosome 1 is affected by susceptibility. The cause of these problems in visualization is the anatomic slant structure of nigrosome 1; each anatomic slant structure individually has an effect on visualization.

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