Association between increased magnetic susceptibility of deep gray matter nuclei and decreased motor function in healthy adults

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A B S T R A C T

In the human brain, iron is more prevalent in gray matter than in white matter, and deep gray matter structures, particularly the globus pallidus, putamen, caudate nucleus, substantia nigra, red nucleus, and dentate nucleus, exhibit especially high iron content. Abnormally elevated iron levels have been found in various neurodegenerative diseases. Additionally, iron overload and related neurodegeneration may also occur during aging, but the functional consequences are not clear. In this study, we explored the correlation between magnetic susceptibility—a surrogate marker of brain iron—of these gray matter structures with behavioral measures of motor and cognitive abilities, in 132 healthy adults aged 40–83 years. Latent variables corresponding to manual dexterity and executive functions were obtained using factor analysis. The factor scores for manual dexterity declined significantly with increasing age. Independent of gender, age, and global cognitive function, increasing magnetic susceptibility in the globus pallidus and red nuclei was associated with decreasing manual dexterity. This finding suggests the potential value of magnetic susceptibility, a non-invasive quantitative imaging measure of iron, for the study of iron-related brain function changes.

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Introduction

Iron is essential for normal functioning of the brain, because it is a co-factor for many enzymes involved in oxygen transport, neurotransmitter synthesis, electron transfer, and myelin production (Moos and Morgan, 2004; Zecca et al., 2004). Plasma iron is not directly available for neuronal cells, but is moved across the blood brain barrier to the neuronal cells through specific transport mechanisms (Burdo and Connor, 2003). Brain tissues store the iron in the forms of ferritin or hemosiderin. Iron and iron storage proteins such as ferritin are found not only in oligodendrocytes but also in microglial cells in the brain. The rates of iron uptake and storage vary significantly across brain regions (Hallgren and Sourander, 1958; Haacke et al., 2005). Specifically, a number of deep brain gray matter structures, including putamen (PU), globus pallidus (GP), caudate nuclei (CN), red nuclei (RN), substantia nigra (SN), dentate nuclei (DN), and others, have much higher iron contents than surrounding tissues (Hallgren and Sourander, 1958). Despite the many important physiological functions of iron, excessive iron can stimulate oxidative damage through the generation of toxic free radicals (Emerit et al., 2001). There is increasing evidence that excessive iron accumulation, especially in the iron rich gray matter regions, is closely involved in mechanisms associated with neurodegeneration in aging, such as Parkinson’s disease and Alzheimer’s disease (Zecca et al., 2004).

Deposition of paramagnetic non-heme iron in brain tissue can lead to a reduction of T2* relaxation rate and altered phase contrast that is measurable by gradient echo (GRE) MRI. While the reduction in T2* has long been used as a measure of brain iron (Gelman et al., 1999; Aquino et al., 2009), the recently developed technique of quantitative susceptibility mapping (QSM) allows the quantification of tissue magnetic susceptibility from GRE signal phase (Liu et al., 2009; Shmueli et al., 2009; de Rochefort et al., 2010; Wharton et al., 2010; Li et al., 2011; Schweser et al., 2011; Wu et al., 2012). The resultant tissue magnetic susceptibility provides a novel contrast with high sensitivity to brain iron. Similar to R2* (1/T2*), accumulating evidence suggests that the magnetic susceptibility of gray matter is linearly proportional to the iron content (Shmueli et al., 2009; Schweser et al., 2011; Bilgic et al., 2012; Langkammer et al., 2012; Wu et al., 2012). Magnetic...
susceptibility is a basic physical property of matter, independent of magnetic field strength, whereas $R_2^*$ scales linearly with field strength (Yao et al., 2009). In addition, magnetic susceptibility mapping typically provides a higher contrast-to-noise ratio and clearer gray–white matter tissue boundary (Rauscher et al., 2005; Li et al., 2011; Deistung et al., 2013; Li et al., 2014b). Thus, QSM is a promising candidate for noninvasive assessment of brain iron.

Early histochemical work revealed that brain iron increases with age (Hallgren and Sourander, 1958). A recent study evaluated the evolution of magnetic susceptibility as a function of age in a cohort of 191 healthy participants from 1 to 83 years of age (Li et al., 2014b). The results indicated that susceptibility of iron-rich, deep gray matter follows an exponential growth trajectory, which agrees well with the known characteristics of brain iron accumulation. Interestingly, MR susceptibilities of these nuclei also exhibited a wider variation of iron content at older ages (e.g., >40 years of age). While existing studies have so far demonstrated the anatomical variations and age-related evolution of magnetic susceptibility, it is unclear if these variations in susceptibility have any functional significance.

Given the role of iron overload in tissue degeneration and the association of deep gray matter in motor function, we hypothesized that in healthy adults, increased magnetic susceptibility of the iron-rich, deep gray matter nuclei is correlated with decline of motor functions, independently of age. In this study, we examined the relation between susceptibility and behavioral performance measures in a cohort of 132 healthy, community-dwelling adults between 40 and 83 years of age. The mean magnetic susceptibility of six subcortical gray matter regions was measured. Participants performed a series of behavioral tests targeting various cognitive and motor abilities. We performed factor analysis to identify the latent variables of the test scores, and correlated the factor scores with the magnetic susceptibility of the six deep gray matter regions.

Materials and methods

Participants

The study cohort was from the Austrian Stroke Prevention Family Study (ASPS-Fam), a prospective, single-center, community-based study on the cerebral effects of vascular risk factors in healthy older adults of the city of Graz, Austria. The ASPS-Fam represents an extension of the Austrian Stroke Prevention Study (ASPS) which was established in 1991 (Schmidt et al., 1999; Schmidt et al., 2003; Schmidt et al., 2005). Between 2006 and 2013, study participants of the ASPS and their first-grade relative were invited to enter ASPS-Fam. Inclusion criteria were no history of previous stroke or dementia and a normal neurologic examination. A total of 381 individuals were included. The entire cohort underwent an extended diagnostic work-up including clinical history, blood tests (including blood iron levels), cognitive testing and a thorough vascular risk factor assessment. All individuals underwent MRI, except for 26 individuals who had contraindications. Magnetic susceptibility imaging was available in a total of 135 participants 40–83 years of age at the time of this study. Three participants were excluded due to the following reasons: (1) missing data from several behavioral tests; (2) obvious signs of bleeding in GP; and (3) poor image quality. Therefore, the total number of participants reduced to 132 ($M = 64.50$, $SD = 10.64$), with mean Mini-Mental State Exam (MMSE) score (Folstein et al., 1975) of 28.19 ($SD = 1.46$). Women were 59% ($n = 78$) of the sample. The study protocol was approved and accepted by the ethics committee of the Medical University of Graz, Austria, and an informed consent was obtained from all participants.

Brain imaging

Each participant was scanned on a 3T scanner with a 12-channel receive array coil at the Medical University of Graz. A spoiled 3D multi-echo gradient-echo sequence (FLASH) was used for quantitative susceptibility mapping. The scan parameters were: in-plane resolution = 0.9 × 0.9 mm², matrix = 256 × 208, flip angle = 20°, TE of first echo = 4.92 ms, echo spacing = 4.92 ms, and number of echoes = 6. The slice thickness was either 4 mm with TR = 68 ms or 2 mm with TR = 35 ms, respectively. The entire brain was covered by the FLASH sequence. Previous studies confirmed that 2-mm and 4-mm slice thicknesses do not produce significantly different magnetic susceptibility values of the iron rich deep gray matter structures (Li et al., 2013b).

To determine the nonlinear warping matrix between the individual participant space and the MNI space, T1-weighted images of the same participants were acquired using a 3D MP-RAGE sequence. The images were acquired in the sagittal view with the following parameters: data matrix = $224 \times 256 \times 176$, 1 mm isotropic resolution, flip angle = 9°, TI = 900 ms, TE = 2.19 ms and TR = 1900 ms.

Quantitative susceptibility mapping

We performed quantitative susceptibility mapping as described previously (Li et al., 2011) under the software STI Suite (Duke University) (Li et al., 2014a). Briefly, the brain was extracted from the magnitude using the brain extraction tool in FSL (Smith, 2002). Phase maps were unwrapped using a Laplacian-based phase unwrapping method (Li et al., 2011). The unwrapped phase maps from all echoes were then normalized by the corresponding echo times to yield the frequency shift using the following equation:

$$
\Delta f = \frac{\sum_{e=1}^{ne} \psi_e}{\sum_{e=1}^{ne} T E_e} \quad [1]
$$

Here, $ne$ is the number of echoes; $\psi$ is the image phase and $\Delta f$ is the frequency shift. This equation assumes linear evolution of phase contrast and assigns different weights for different echoes according to their TEs. This approach is valid for the evaluation of magnetic susceptibility of gray matter structures, with the advantage of higher signal-to-noise level than susceptibility mapping using only a single echo. The background frequency was removed using a variable-filter-radius SHARP method (Schweser et al., 2011). Specifically, the diameter of the spherical mean filter decreases from a maximum value of 25 mm towards 1 mm at the brain boundary (Liet al., 2011; Wu et al., 2012). Susceptibility maps were then derived from the brain tissue frequency shift using the LSQR method (Li et al., 2011).

Regions of interest

We obtained the deep gray matter regions of interest (ROIs) by warping a customized common atlas created in the MNI space to each individual participant space using FSL (FMRIB, University of Oxford, UK). A schematic diagram describing this procedure is shown in Fig. 1. Briefly, we registered the T1-weighted images to the standard template (MNI152_T1_1mm) using FNIRT. We then used the resulting registration matrices to wrap the susceptibility maps to the MNI space and then averaged the maps to generate mean susceptibility. We initially created the atlas for the deep gray matter nuclei using the Harvard–Oxford subcortical atlas and Talairach atlas in FSL; we then manually refined them based on the structural boundaries shown in the mean susceptibility maps.

Examples of the six ROIs, and their associated mean magnetic susceptibility values, are presented in Fig. 2. Six major deep gray matter nuclei were segmented, specifically the PU, GP, CN, RN, SN, and DN (Fig. 2, Panels A–C). This study-specific common atlas for deep gray matter nuclei was then warped to each individual participant space using FSL. The ROIs of each participant were visually examined and minor refinement of the ROIs was made when necessary. As illustrated in Fig. 2, Panels D and E, the ROIs varied in size as well as the mean level of magnetic
susceptibility. Thus, as noted in the Results section, the susceptibility values were adjusted statistically for ROI size.

Behavioral tests

Each participant completed a series of behavioral tests: perceptual motor functioning, verbal working memory, executive functioning, and verbal fluency within one month of MRI scan. These tests yielded a total of 11 dependent variables, as described as follows.

Purdue-pegboard-test (Lafayette Instrument Company, USA)

This test is a clinical measure used to assess manual dexterity and perceptual-motor speed (Tiffin and Asher, 1948). The pegboard features two parallel lines of holes running vertically down the center of

Fig. 1. A schematic representation of the image analysis pipeline (A) and the mean magnetic susceptibility maps in different axial locations (B).

Fig. 2. Regions of interest (ROIs) and their corresponding magnetic susceptibility values. A–C: three axial slices of quantitative susceptibility maps. D: Mean susceptibility in ppm for each ROI, ±1 SE; E: Mean size in mL for each ROI, ±1 SE. PU = putamen; GP = globus pallidus; RN = red nuclei; SN = substantia nigra; DN = dentate nuclei; CN = caudate nuclei.
the board. This test consisted of four subtests. In the first portion of
the test, a participant was given 30 s to place as many pins in the
holes as possible using the right hand. In the second portion of
the test, this process was repeated using the left hand. In the third portion of
the test, the participant had another 30 s to place pins into the
holes using both hands simultaneously. In the final portion of
the test, usually called the assembly test, the participant must as-
semble pins, washer, and collar in a fixed order using the left and
the right hand alternatively. The participant had 60 s to complete
as many assemblies as possible. Thus, the Purdue pegboard test
yielded four dependent variables: a) pins/30 s right; b) pins/30 s
left; c) pins/30 s both; d) assembles/60 s assembly.

**Digit span test**

This subtest of the Wechsler Adult Intelligence Scale IV is composed
of digits-forward and digits-backward tasks, and is used to measure ver-
bal working memory (Wechsler, 2003). In these tasks, participants lis-
tened to a series of discrete digits and subsequently recalled the series
of digits in the same (i.e., digits-forward task) or reverse
(i.e., backward digit span task) order of presentation. Each task com-
prised eight difficulty levels with two trials for each level. The test
started with a two-digit sequence, and the series of digits were subse-
quent increased by one digit every two trials. The task was
discontinued if the participant failed to recall two consecutive
series with the same number of digits. Each correct trial is awarded a score of
1, with a total maximum score of 16 points. The digit span test yielded
two measures of memory span: digits-forward score and digits-
backward score.

**Wisconsin Card Sorting Test (WCST, Heaton et al., 1993)**

This test was designed to measure the ability to display flexibility
in the face of changing rules. It is assumed that successful completion
of this test requires engagement of executive functions (Jodzio and
Biechowska, 2010). During this task, four stimulus cards with shapes
on them were presented to the participant. The stimulus cards dif-
f ered in color, number, and the form of the shapes. The participant
was asked to match a series of cards to one of the four stimulus
cards based on one of the rules (color, number or form of shape).
The correct rule was not revealed to the participant. Instead, the par-
ticipant was instructed to determine the rule based on the feedback
(right or wrong) he/she received upon response and was told that
the rule might change without notice. The rule actually changed
every time the participant completed ten consecutive correct trials.
The WCST yielded three dependent variables: 1) the number of cat-
ergories completed, 2) the number of perseverative errors
(i.e., errors due to a failure to switch from the previous rule), and
3) the sum of non-perseverative and perseverative errors.

**Trail making test B (War Department Adjutant General’s Office, 1944; Reitan and Wolfson, 1985)**

This test measures executive functioning, psychomotor speed, and
visual scanning. Participants were instructed to draw lines to connect
25 circled numbers and letters in an alternating numeric and alphabetic
sequence (i.e., 1-A-2-B) as rapidly as possible. The dependent variable
was the time in seconds to finish the test.

**Semantic fluency test**

To measure semantic fluency, we tabulated the number of super-
market items the participants named in 1 min (“name things you can
buy in a large supermarket in 1 min”) (Lezak, 1995). The dependent
variable was the number of items named over 1 min.

Mean values of the behavioral tests, and clinical measures of blood
iron levels, are presented in Table 1.

### Results

**Magnetic susceptibility**

Because the ROIs varied widely in size (i.e. the volumes of the deep
brain nuclei) (Fig. 2), all analyses of the magnetic susceptibility values
were adjusted for ROI size in the following manner. For each ROI, we
created a regression model in which ROI size was a predictor of suscept-
ability. We then used the residual of this model, for each participant,
for use in further analyses. Because the residuals are by definition uncor-
related with size, this method created a size-independent measure of sus-
cceptibility for each ROI. We then formed standardized (z) scores of
these size-residualized susceptibility values for each ROI, so that the
ROI susceptibility values would have a common scale. Further, prelimi-
inary analyses indicated no reliable effects of gender, but gender was
added as a covariate to all analyses.

We have previously shown that the susceptibility of several deep
grey matter nuclei evolves exponentially as a function of age (Li et al.,
2013b). Susceptibility typically increases nonlinearly at young ages
(1–20 years), then either approaching a plateau or increasing linearly
at older ages (>40 years). Across these six ROIs, an age-related increase
in magnetic susceptibility was evident within the putamen (Fig. 3, Panel A),
but the effect of age was not significant within any of the other ROIs.
Further analyses of the susceptibility values included age as a covariate.

**Relation between behavioral performance and magnetic susceptibility**

Prior to analysis, all demographic and behavioral variables were
transformed to z scores based on the sample mean and standard devia-
tion per variable. We performed an exploratory factor analysis of the 11
behavioral measures to identify the independent, latent variables of be-
havioral performance. We then used the factor scores representing
these latent variables as dependent variables in regression analyses
with participants’ years of age and size-residualized susceptibility from
the six ROIs as predictors.

Results of the factor analysis revealed that the eigenvalues of the first
two factors were greater than 1.0, and these factors accounted for 84% of
the common variance. Following Varimax rotation, we used 0.60 as a
threshold for significant factor loadings (Table 2), on the basis of sample
size and number of variables (Buja and Eyuboglu, 1992). The first factor
comprised all four variables from the Purdue pegboard test, and we in-
terpret this factor as representing manual dexterity. The second factor
comprised two variables from the Wisconsin Card Sorting Test, and we in-
terpret this factor as representing executive function. We conduct-
cred correlations between the factor scores and age, partialled for the ef-
fects of gender and global cognitive functioning (MMSE). The first
factor representing manual dexterity exhibited significant decline as a
function of increasing age (Fig. 4, Panel A), whereas the second factor

<table>
<thead>
<tr>
<th>Variable</th>
<th>M</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Purdue pegboard: right (number completed)</td>
<td>12.58</td>
<td>2.46</td>
</tr>
<tr>
<td>Purdue pegboard: left (number completed)</td>
<td>12.30</td>
<td>2.27</td>
</tr>
<tr>
<td>Purdue pegboard: left + right (number completed)</td>
<td>10.29</td>
<td>2.21</td>
</tr>
<tr>
<td>Purdue pegboard: assembly (number completed)</td>
<td>24.89</td>
<td>7.31</td>
</tr>
<tr>
<td>Digit span: forward (items)</td>
<td>6.20</td>
<td>1.14</td>
</tr>
<tr>
<td>Digit span: backward (items)</td>
<td>4.25</td>
<td>1.32</td>
</tr>
<tr>
<td>WCST: errors</td>
<td>18.39</td>
<td>12.25</td>
</tr>
<tr>
<td>WCST: perseverative responses</td>
<td>7.04</td>
<td>4.53</td>
</tr>
<tr>
<td>WCST: categories</td>
<td>5.66</td>
<td>1.00</td>
</tr>
<tr>
<td>Trials B (sec)</td>
<td>115.10</td>
<td>58.90</td>
</tr>
<tr>
<td>Word fluency (items/min)</td>
<td>23.99</td>
<td>6.93</td>
</tr>
<tr>
<td>Iron (ug/dL)</td>
<td>105.48</td>
<td>32.04</td>
</tr>
<tr>
<td>Transferrin (mg/dL)</td>
<td>252.74</td>
<td>44.85</td>
</tr>
<tr>
<td>Ferritin (ug/L)</td>
<td>130.41</td>
<td>109.96</td>
</tr>
</tbody>
</table>

Note: n = 132. WCST = Wisconsin Card Sorting Test.
representing executive function did not exhibit significant age-related decline (Fig. 4, Panel B).

The bivariate correlations between susceptibility and the behavioral factor scores, for each ROI considered independently, are presented in Table 3. These correlations are partialed for the effects of age, gender, and global cognitive function (MMSE). For manual dexterity, a decreasing behavioral score was associated with increasing magnetic susceptibility in both the GP and RN (Fig. 5). Given the relation between magnetic susceptibility and manual dexterity for these two ROIs, we explored potential age-related differences in this pattern. We split the sample into two groups at the median age value (67.4 years) and performed bivariate susceptibility–dexterity correlations within each sub-group, partialed for age, gender, and MMSE. For the younger group \((n = 67)\), the susceptibility–dexterity correlation was significant for the GP, \(r = -0.318, p < 0.01\), but not for the RN, \(r = -0.137, p = 0.28\), whereas for the older group \((n = 65)\), the correlation was significant for the RN, \(r = -0.362, p < 0.01\).

**Table 3**
Rotated factor loadings for factors 1 and 2 from an exploratory factor analysis of 11 behavioral tests.

<table>
<thead>
<tr>
<th>Behavioral Tests</th>
<th>Factor 1 (Manual dexterity)</th>
<th>Factor 2 (Executive function)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Purdue right</td>
<td>0.768</td>
<td>0.112</td>
</tr>
<tr>
<td>Purdue left</td>
<td>0.864</td>
<td>-0.002</td>
</tr>
<tr>
<td>Purdue both</td>
<td>0.819</td>
<td>0.018</td>
</tr>
<tr>
<td>Purdue assembly</td>
<td>0.766</td>
<td>0.068</td>
</tr>
<tr>
<td>Digit symbol forward</td>
<td>0.187</td>
<td>0.084</td>
</tr>
<tr>
<td>Digit symbol backward</td>
<td>0.092</td>
<td>0.086</td>
</tr>
<tr>
<td>WCST error</td>
<td>-0.083</td>
<td>-0.682</td>
</tr>
<tr>
<td>WCST category</td>
<td>0.052</td>
<td>0.854</td>
</tr>
<tr>
<td>WCST perseverative error</td>
<td>0.019</td>
<td>-0.291</td>
</tr>
<tr>
<td>Trail making</td>
<td>-0.422</td>
<td>-0.221</td>
</tr>
<tr>
<td>Semantic fluency</td>
<td>0.340</td>
<td>0.007</td>
</tr>
</tbody>
</table>

Note: \(n = 132\). Factor loadings \(>0.60\) are presented in bold. WCST = Wisconsin Card Sorting Test.

Fig. 3. Magnetic susceptibility values as a function of age, for each region of interest. The susceptibility values are standardized (z) scores, residualized for region size. Correlations are partialed for gender. A significant age-related increase in susceptibility was evident only within the putamen.
Within the GP was associated with decreasing manual dexterity (Fig. 5, predictor, independent of the other ROIs. Increasing magnetic susceptibility (executive function), the overall model with all predictors was not significant, indicating that the variation in the correlations across the age sub-groups was not reliable.

We further obtained an estimate of age-related differences in region size by standardizing each ROI size in mL to the sample mean and standard deviation and then averaging these standard (Z) scores across participants. The average ROI size decreased as a function of increasing age, \( r = -0.262, p < 0.05 \), but not for the GP, \( r = -0.205, p = 0.11 \). For each ROI, however, direct comparison of the correlations for the two subgroups (with Fisher’s r to z transformation), was not significant, indicating that the variation in the correlations across the age sub-groups was not reliable.

The magnetic susceptibility values were all correlated positively across the ROIs, with bivariate correlations ranging from \( r = +0.197, p < 0.05 \), to \( r = +0.571, p < 0.001 \), across the 15 pairs of ROIs. We thus conducted two linear regression analyses, one for each of the latent variables from the factor analysis, representing manual dexterity and executive function, with all of the ROI values entered simultaneously, thus controlling the shared variance among the ROIs. Each regression model contained nine predictor variables: the six susceptibility values from the six ROIs, plus three demographic variables (gender, age, and MMSE) to control for individual differences theoretically unrelated to susceptibility and the behavioral outcome measure.

In the regression analysis of the first behavioral factor (manual dexterity), the nine predictors combined accounted for 37% of the variance in manual dexterity, \( F(9, 122) = 7.82, p < 0.0001 \) (Table 4). Age was a significant predictor, reflecting the age-related decline in this measure. Among the six susceptibility variables, only the GP was a significant predictor, independent of the other ROIs. Increasing magnetic susceptibility within the GP was associated with decreasing manual dexterity (Fig. 5, Panel A). In the regression analysis of the second behavioral factor (executive function), the overall model with all predictors was not significant, \( F(9, 122) < 1.0 \), and none of the individual predictors was significant (Table 5).

Correlations conducted between the three biomarkers of peripheral blood iron and the susceptibility values did not yield any significant effects (Table 6).

**Discussion**

In this study, we investigated the relationship between magnetic susceptibility of iron rich deep gray matter and brain functions measured by behavioral tests. Factor analysis of the behavioral tests identified two latent variables: manual dexterity and executive function. The manual dexterity factor was more sensitive than executive function to the effects of magnetic susceptibility. We found that, controlling for individual differences in gender, age, and global cognitive function, decreased manual dexterity was significantly correlated with increased magnetic susceptibility of GP and RN, but not the susceptibility of other gray matter structures (Table 3 and Fig. 5). When controlling for the shared variance among the ROIs, the age-related effect remained significant for the GP, but not for the RN (Tables 4 and 5), indicating a stronger effect for the GP. As the GP is the gray matter structure containing the highest amount of non-heme iron (Fig. 2, Panel D), these results suggest that high iron contents in GP can predispose individuals to higher risk of iron-overload induced motor function decline.

While the six different gray matter nuclei share a common feature of high iron content, they belong to different structural and functional units of the brain. The CN, PU, GP and SN are the main components of the basal ganglia, which are known to play major roles in regulating and modulating motor control functions through the widespread interconnections with the neocortex and the prefrontal and premotor cortices (Kandel and Schwartz, 1985). Consistent with this knowledge, we observed significant correlations between the manual dexterity factor score and the magnetic susceptibility of GP. These results suggest that iron overload in GP is associated with a negative impact on brain motor functions. Overall, our results are in good agreement with a previous exploratory study using 10 participants indicating that the field-dependent R2 increase (FDRI) values, another measure of brain iron, of basal ganglia nuclei also correlated with motor performance (Sullivan et al., 2009). It is worth noting that our results did not reveal a significant correlation between the magnetic susceptibility of SN and motor function scores, although it is well known that the motor function decline in Parkinson’s disease is associated with dopamine cell loss and increased iron content in SN. A previous study also documented the increased magnetic susceptibility in Parkinson’s disease (Lotfi Pour et al., 2012). It is possible that the iron accumulation in SN during normal aging in our study group did not reach the extent as found in pathological tissue damage.

**Table 3** Correlations of magnetic susceptibility with manual dexterity and executive function.

<table>
<thead>
<tr>
<th>Region</th>
<th>Manual dexterity</th>
<th>Executive function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Putamen</td>
<td>-0.058</td>
<td>-0.047</td>
</tr>
<tr>
<td>Globus pallidus</td>
<td>-0.216**</td>
<td>-0.077</td>
</tr>
<tr>
<td>Red nuclei</td>
<td>-0.199*</td>
<td>-0.120</td>
</tr>
<tr>
<td>Substantia nigra</td>
<td>-0.074</td>
<td>-0.099</td>
</tr>
<tr>
<td>Dentate nuclei</td>
<td>-0.062</td>
<td>-0.158</td>
</tr>
<tr>
<td>Caudate nuclei</td>
<td>0.028</td>
<td>0.036</td>
</tr>
</tbody>
</table>

Note. \( n = 132 \). Values are Pearson r correlations, partialed for the effects of age, gender, and global cognitive function (MMSE). Magnetic susceptibility values were residualized for the effects region size.

* \( p < 0.05 \).

** \( p < 0.01 \).
The RN is a prominent brain stem structure, which is dorsally located to SN but is involved in different neuronal networks from that of basal ganglia. The origin of RN is suggested to be related to the advent of limbs, or limb-like structures during evolution. The RN plays important roles in motor control functions in lower vertebrates, but its importance related across the ROIs. As a result, the selection of ROIs and the participant inclusion may influence the results of the statistical analysis. In this study, the ROIs were created and refined before statistical analysis. All 132 participants were included in the analysis as all are clinically considered healthy. Although global cognitive function as represented by MMSE was controlled statistically, 14 participants had MMSE scores < 27, and further research should investigate whether the relation between magnetic susceptibility and behavioral performance varies in relation to cognitive status. Another limitation is that the behavioral factors in Fig. 4 are specific to individual tests. All the variables comprising the first factor came from Purdue pegboard, and the second factor comprised only two variables, both of which came from WCST. Thus, the factors do not represent general cognitive abilities beyond these individual tests. Further, Purdue pegboard contributed more variables to the factor analysis than the other tests, which may have influenced its role as the primary factor. Thus, studies with more behavioral tests and more subjects may help us further investigate and confirm our findings.

Our results also showed that magnetic susceptibilities of the iron rich nuclei are not correlated with blood iron, transferrin or ferritin (Table 6). These results are reasonable given the multiple regulating factors during iron transport, uptake and storage. As such, neuroimaging methods, such as QSM, can offer more accurate means to investigate the effect of iron overload on the neurological system than the peripheral blood measurements. As iron overload has been suggested to be involved in mechanisms underlying major neurological disorders, such as Alzheimer’s disease, Parkinson’s disease, and Huntington’s disease (Kandel and Schwartz, 1985; Zecca et al., 2004), this type of correlation thus indicate that the negative correlation between susceptibility of GP and motor function is likely due to the accumulation of iron.

A limitation of this study is that magnetic susceptibility is highly correlated across the ROIs. As a result, the selection of ROIs and the participant inclusion may influence the results of the statistical analysis. In this study, the ROIs were created and refined before statistical analysis. Although increased magnetic susceptibility in the RN did exhibit a significant relation to decreased manual dexterity, the RN-manual dexterity relation was not statistically independent of the stronger behavioral correlation exhibited by the GP.

Magnetic susceptibility of gray matter nuclei is predominantly determined by the level of iron concentration. Our finding that susceptibility of GP is negatively correlated with motor function in healthy adults is consistent with prior findings that iron accumulates in specific gray matter regions and that iron overload is associated with functional decline during aging and in neurological diseases (Zecca et al., 2004). Another study also reported that a calorie-restricted diet can decrease brain iron accumulation and preserve motor performance (Kastman et al., 2010). While the biological mechanism of age-related iron accumulation is unknown, the negative correlation between susceptibility and motor function is not simply due to age, as susceptibility of GP was not significantly correlated with age in our cohort, and our analyses statistically controlled for age-related effects in all ROIs. Further, our analyses controlled statistically for ROI size in all analyses. Our findings therefore indicate that the negative correlation between susceptibility of GP and motor function is likely due to the accumulation of iron.
studies would open a new avenue to investigate the roles of iron accumulation during neurodegeneration.

In conclusion, we found that the manual dexterity, as measured by the latent variables of behavioral tests, is negatively correlated with magnetic susceptibility in the GP. As GP contains the highest amount of non-heme iron in the brain, this result suggests that magnetic susceptibility of the GP is a potential marker of motor function decline. Our results also support the added value of QSM in studying the impact of iron deposition on brain function compared to simply using the levels of peripheral blood iron or iron-related transport or storage proteins.

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