Brain structure in healthy adults is related to serum transferrin and the H63D HFE polymorphism


Control of iron homeostasis is essential for healthy central nervous system function: iron deficiency is associated with cognitive impairment, yet iron overload is thought to promote neurodegenerative diseases. Specific genetic markers have been previously identified that influence levels of transferrin, the protein that transports iron throughout the body, in the blood and brain. Here, we discovered that transferrin levels are related to detectable differences in the macro- and microstructure of the living brain. We collected brain MRI scans from 615 healthy young adult twins and siblings, of whom 574 were also scanned with diffusion tensor imaging at 4 T. Fiber integrity was assessed by using the diffusion tensor imaging-based measure of fractional anisotropy. In bivariate genetic models based on monozygotic and dizygotic twins, we discovered that partially overlapping additive genetic factors influenced transferrin levels and brain microstructure. We also examined common variants in genes associated with transferrin levels, TF and HFE, and found that a commonly carried polymorphism (H63D at rs1799945) in the hemochromatotic HFE gene was associated with white matter fiber integrity. This gene has a well-documented association with iron overload. Our statistical maps reveal previously unknown influences of the same gene on brain microstructure and transferrin levels. This discovery may shed light on the neural mechanisms by which iron affects cognition, neurodevelopment, and neurodegeneration.

Iron and the proteins that transport it are critically important for brain function. Iron deficiency (ID) is the most common nutritional deficiency worldwide (1). Iron-deficient diets lead to poorer cognitive achievement in school-aged children (2). In rural areas where ID anemia is prevalent, iron supplements can increase motor and language capabilities in children (3). ID also impairs dopamine metabolism in the brain, particularly in the caudate and putamen regions (4). ID clearly has adverse effects on cognitive development, but iron overload in later life is also associated with damage to the brain. Brain iron regulation is disrupted in several neurodegenerative diseases. Neuroimaging methods reveal abnormally high brain iron concentrations in Alzheimer’s disease (5), Parkinson disease (6), and Huntington disease (7). High iron concentrations may cause neuronal death (8, 9).

As deficiency and excess of iron can negatively impact brain function, the regulation of iron transport to the brain is crucial for cognition. Iron is transported throughout the body by the iron-binding protein transferrin. The interaction between transferrin and the transferrin receptors appears to regulate iron transport (10). When iron levels are low, the liver produces more transferrin for increased iron transport. In humans, transferrin can increase in iron-deficient states, which may help to distinguish ID anemia from anemia of chronic disease (11). Dietary ID has also been shown in rats to elevate the concentration of transferrin in the brain (12), specifically in the hippocampus and striatum (13). Transferrin is also decreased in cases of iron overload (14).

The gold standard for determining accurate iron measures is obtained from invasive bone marrow or liver tests, which are impractical for general applications. Serum levels of iron fluctuate greatly (15) and depend on dietary factors such as vitamin C intake (16) and the time of blood collection (17). Transferrin is arguably a more reliable and reproducible index of the long-term availability of iron to the brain (18, 19). In fact, in a 2-y study of postmenopausal women (20), total iron-binding capacity (equivalent to transferrin concentration) was a more reliable measure of iron status [0.60; 95% confidence interval (CI), 0.44–0.76], whereas serum iron measures varied more (0.50; 95% CI, 0.22–0.65). Transferrin is therefore used as a more reproducible measure to infer iron availability to the neural pathways.

As iron is a key determinant of neural development and degeneration, we set out to investigate whether brain structure in healthy adults depends on serum transferrin levels. We scanned 615 young adult twins and siblings with standard MRI. A total of 574 of them were also scanned with diffusion tensor imaging (DTI) to assess volumetric and microstructural white matter differences potentially associated with variations in serum transferrin levels measured during adolescence.

The participants in our study were healthy young adults, in whom iron overload is unlikely. We instead expected iron levels toward the lower end of the normal range might lead to a poorer developmental phenotype in the brain of these young adults. The brain synthesizes transferrin itself, so serum transferrin is not necessarily indicative of the levels of brain transferrin. However, in healthy populations without iron overload or hemochromatosis, all iron in the plasma is bound to transferrin (10). Iron enters the brain primarily by transport through the blood–brain barrier (21), yet transport through the blood–CSF and cellular–plasmalemna barriers have also been described (10). Upon binding to its receptor, transferrin is thought to be mostly released back into the blood stream (although some transcytosis of transferrin may occur). The iron can then bind to transferrin synthesized in the oligodendrocytes of white matter (22, 23).

This article is a PNAS Direct Submission.

Author contributions: N.J., O.K., and P.M.T. designed research; N.J. and P.M.T. performed research; K.L.M., G.L.d.Z., S.E.M., G.W.M., J.B.W., N.G.M., M.J.W., and A.W.T. contributed new reagents/analytic tools; N.J., O.K., D.P.H., J.L.S., and P.M.T. analyzed data; and N.J. and P.M.T. wrote the paper.

The authors declare no conflict of interest.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1105543109/-/DCSupplemental.
Most of the brain’s iron is found in oligodendrocytes, where it supports myelination (24). Oligodendrocytes also maintain iron homeostasis in the brain. Our primary hypothesis was that we might find poorer white matter integrity in adulthood in those who had lower iron levels available during development, as high transferrin level is often a sign of the liver reacting to lower iron availability. We therefore framed our hypothesis by testing if serum transferrin levels in adolescence were related to fractional anisotropy (FA; measured later in adulthood from DTI scans of the brain). Lower FA can be a sign of less mature or poorer myelination.

We further hypothesized that brain structure volumes in iron-rich regions might be lower in people with high serum transferrin levels. Iron levels are highest in basal ganglia and substantia nigra (25). By measuring brain volumes regionally with tensor-based morphometry (TBM), we predicted that we might find insufficiently developed (i.e., smaller) subcortical structures in those with higher transferrin levels. ID alters dopamine metabolism in the caudate and putamen (4), so we predicted that people with high transferrin (and, by implication, lower brain iron) might have lower volumes for dopamine-containing structures, such as the caudate. Finally, we expected lower hippocampal volumes, as iron-deficient rats have lower iron concentrations in the hippocampus (13), a region vulnerable to neuronal loss in neurodegenerative disease (26).

Genetic factors explain 66% and 49% of the variance in serum transferrin levels in men and women, respectively (17). As such, if transferrin is found to be associated with neuroanatomical differences, we might expect that common genes influence both brain structure and transferrin levels. To understand such shared genetic contributions to brain variation and transferrin, we used a twin design. Many neuroimaging studies of identical and fraternal twins reveal substantial genetic contributions to brain structure (27–30) and function (31, 32). Cross-twin cross-trait designs can also discover overlapping (i.e., pleiotropic) genetic influences on very different biological traits, such as brain volume (33) and fiber integrity and IQ (34).

After discovering a common genetic basis for transferrin levels and brain fiber integrity, we hypothesized that genes modulating transferrin also play a role in brain structure within the same regions. We performed exploratory tests on all SNPs within the two major transferrin related genes: the transferrin gene, TF, on chromosome 3, and in the HFE gene on chromosome 6, where a handful of SNPs have been found to explain a remarkable 40% of the genetic variance in serum transferrin levels (35). We performed exploratory tests on all these SNPs and additional imputed ones (to HapMap2) within the same genes.*

Genes influencing transferrin are not the only cause of variation in iron levels measured in the blood serum. However, they do influence the limited amount of serum iron that becomes transported into the brain. Therefore, we expected genes that influence transferrin levels to show associations with brain structure. Some variants increase the risk for iron overload late in life, and these may also increase the availability of brain iron for developmental processes such as myelination. If high iron levels improve myelination, we might expect to see increased fiber integrity as measured through DTI.

**Results**

Serum transferrin levels for the 615 individuals in our study ranged from 1.89 to 5.18 g/L of serum (mean, 2.99 ± 0.37 g/L; median, 2.96 g/L). Regressions fitted to data at each voxel in the brain DTI and MRI scans revealed significant associations with microstructural variations in diffusion anisotropy (Fig. 1). There were also strong associations between transferrin levels and gross anatomical volume differences (Fig. 2), even after controlling for age and sex. As ID is known to reduce myelination, we had expected, in advance, to find a negative association between transferrin and FA (36); this was in fact observed.

Transferrin Levels Relate to Neuroanatomical Structure. As noted, cross-twin cross-trait models can determine whether a partially overlapping set of genes contributes to two traits of interest, such as fiber integrity (assessed by using FA) and transferrin levels. If this is the case, common genetic influences mediate the observed correlation between the two measures. Before examining the full cross-twin cross-trait model, we independently assessed these correlations within each group of twins; if the variables correlate more strongly for monozygotic (MZ) than for dizygotic (DZ) twin pairs, then we can infer the greater difference is a result of additive genetic factors. Fig. 3 shows the correlations between FA and transferrin levels in MZ twins and DZ twins separately, highlighting the higher magnitude of the correlations for MZ than DZ pairs.

Cross-Twin Cross-Trait Analysis of Shared Genetic Determination. We performed cross-twin cross-trait heritability analysis starting from the full bivariate model as described in Methods; we remove individual components one by one to determine the best fitting model. For both MRI- and DTI-based bivariate ACE models, the AE model fitted the best for transferrin and the full ACE model fitted best for the imaging measures. This means that genetic effects were detected in both cases, and the effects of common rearing environment were also detectable for the imaging measures. The path diagram for the best fitting model is shown in Fig. 4.

The cross-twin cross-trait correlation was then computed from the best fitting model. Significance of the correlation was determined by removing the A component of the path model as described in Methods. The additive genetic determinant of voxel-wise FA measures (Fig. 5) and transferrin showed significant overlap after multiple comparisons correction using the false discovery rate (FDR) procedure (37). Although suggestive, no significant overlap was detected between the additive genetic

```bash
# Table
<table>
<thead>
<tr>
<th>Variable</th>
<th>Std Error</th>
<th>P-Value</th>
<th>FDR Corrected</th>
</tr>
</thead>
<tbody>
<tr>
<td>FA</td>
<td>0.12</td>
<td>&lt;0.01</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Transferrin</td>
<td>0.15</td>
<td>&lt;0.02</td>
<td>&lt;0.05</td>
</tr>
</tbody>
</table>
```

Fig. 1. Voxel-wise associations, between FA, a measure of white matter fiber integrity derived from the DW images, and serum transferrin levels in 574 subjects (five of whom had repeated scans). There are significant associations in the external capsule, superior longitudinal fasciculus, and the cingulum bilaterally. As transferrin levels increase, the diffusivity across the axons also tends to decrease by approximately 0.025 units for every g/L unit increase in the serum transferrin level. Significance was confirmed by enforcing a regional control over the FDR as described by Langers et al. (70) at the 5% level. Corrected P values of association are shown. Maps are adjusted for effects of age and sex; random-effects regression accounted for familial relatedness and the use of repeated scans. Values shown represent the regression coefficient (or slope) of the transferrin level term, after accounting for covariates.
Genetic Associations. After filtering the SNPs in TF and HFE available in our imputed sample by minor allele frequency (MAF) greater than 0.05, 42 SNPs remained. SNPs chosen for analysis are listed in Table S1, along with their MAF according to the CEU population from HapMap. As a result of linkage disequilibrium, the effective number of SNPs tested (38, 39) was 20. When the significant voxels of the cross-twin cross-trait associations were clustered into regions of interest (ROIs), six survived a cluster threshold size of 27 voxels, corresponding to the size of a voxel with all its surrounding neighbors, or a 3 × 3 × 3 cube. These ROIs are shown in Fig. S1. Genetic associations of the 42 (effectively 20) SNPs performed in these six regions revealed a significant association of the HFE rs1799945 SNP (also known as the H63D polymorphism) with the mean FA in the cluster along the left external capsule (P = 0.00017). The results of all of the genetic associations per ROI are also listed in Table S1.

Additionally, in the full sample of 565 genotyped subjects with serum transferrin levels available, we found that the H63D minor allele was associated with decreased transferrin levels as expected (r statistic = 1.801, one-tailed P = 0.0361).
Post Hoc Voxel-Wise Analysis of H63D HFE Missense Polymorphism. In our post hoc analysis, we performed a voxel-wise association of FA with the H63D polymorphism across the entirety of the white matter region. The SNP frequency information for this polymorphism in our sample is available in SI Methods. The map of voxel-wise associations of H63D to FA values was found to be significant in regions including the external capsule, and portions of the genu of the corpus callosum not initially found to have significant transferrin related associations (Fig. 6).

Discussion

ID, iron overload, and abnormalities in iron concentrations localized to particular structures in the brain have been linked to neurodevelopmental and neurodegenerative disorders. Fig. 7 shows a schematic illustration relating several biological processes that motivated this study. Both brain structure and the iron transport protein, transferrin, are under strong genetic control, so we used a twin design to find brain regions with genetic determinants in common with transferrin. We were able to establish some previously unknown links between transferrin levels (and associated genes) and brain structure in 615 healthy young adults.

Our analysis had three main findings. First, serum transferrin levels, measured during adolescence, were associated with both macro- and microneuroanatomical variations in a regionally selective pattern in later adulthood (~9 y after blood was drawn). Second, these associations with white matter integrity were mediated by overlapping sets of genes. This was evident from the cross-twin cross-trait correlations between transferrin levels and white matter anisotropy. Third, we found that the HFE H63D polymorphism, well known for its association to iron overload (40, 41), influences both serum transferrin levels and white matter microstructure in the external capsule. This points to a direct link between blood serum related genomic variation and brain structure (Fig. 7, dashed lines).

Iron is important for neural development early in life. In rat brains, iron and transferrin are at extremely high levels, despite low brain transferrin mRNA levels before closure of the blood–brain barrier (42). Even after the barrier develops, serum transferrin levels, which are under high genetic control, influence how much iron is transported to the brain for crucial processes of development, such as myelination. Here we uncovered an association between brain structure in young adults and serum transferrin levels measured during their adolescent years.

By measuring transferrin levels 8 to 12 y before the imaging study, we were interested in knowing whether iron availability in this developmentally crucial period might impact the organization of the brain later in life. Adolescence is a period of high vulnerability to brain insults, and the brain is still very actively developing (43). Transferrin levels, measured before the brain is fully mature, may be especially relevant for the adult brain. Transferrin levels fall with age in children and adolescents, and older adolescents show similar ranges to adults (15, 44). Children have higher transferrin levels than adults, perhaps in response to physiologically low iron stores. By averaging transferrin levels assessed repeatedly at various ages (12, 14, and 16 y of age), we estimated the iron availability to the brain during adolescence. We relied on previous work showing that transferrin measures are stable and can be reliably collected (19), with high sensitivity and specificity, which makes associations easier to detect.

As key components of white matter, oligodendrocytes—the glial cells that produce myelin to insulate axons—stain for iron more than any other cell in the brain (24); these cells are the primary location for iron in the central nervous system (45). Fig. 1 shows the negative association between serum transferrin levels and the diffusion-based measure of integrity, FA, in various brain regions, including the external capsule, cingulum, and superior longitudinal fasciculus. As FA can represent the degree of fiber integrity, myelination, or coherence in white matter fibers, the direction of this association is in line with previous reports indicating hypomyelination in cases of ID (36). Our findings of transferrin associations in human white matter tracts are consistent with previous studies of brain iron levels in rats. Connor et al. (46), in a histochemical study of iron staining in the developing rat brain, found major foci of iron staining in the cingulum, superior portions of the internal capsule, and the base of the external capsule.

As shown in Fig. 2, regional brain volume deficits are seen bilaterally in the caudates, the third ventricle, and in the temporoparietal white matter as transferrin levels increase. The caudate is particularly important in cognition, learning, and memory (47); as increased transferrin levels have been implicated in ID, the inverse relation between transferrin levels and caudate volume may indicate underdevelopment of caudates in ID, in line with evidence of poorer cognition in children with ID. If severe enough, a caudate volume reduction related to elevated transferrin levels may explain why certain developmental cognitive deficits are associated with ID. This finding fits with our previous hypothesis that insufficient iron transport to the brain may hinder development of subcortical structures.

Although transferrin levels are low, transferrin in the CSF is fully saturated with iron (48); the regional deficit in the volume of the third ventricle is therefore intriguing, as it may indicate an altered pattern and/or rate of transport for transferrin-bound iron.

Fig. 6. Corrected p-map shows the HFE H63D associations with FA voxel-wise throughout the white matter. When regressing on the minor allele, there is a positive association between the number of minor alleles and the FA values. Significance was confirmed by enforcing a regional control over the FDR as described by Langers et al. (70) at the 5% level. We adjusted for effects of age, sex, and total serum iron levels to be consistent with the previous tests. Positive correlations were not significant.

Fig. 7. Several known relationships motivated our study (solid black lines); dashed lines show relationships we wanted to test. Genetic and environmental factors (e.g., diet) affect iron stores in the body; the liver synthesizes more transferrin in response to low iron stores. Our first goal was to relate transferrin levels to brain structure in healthy young adults. Our twin design determined if overlapping sets of genes influence transferrin levels and brain structure, as both are highly heritable. Transferrin levels are genetically modulated mainly through two genes (HFE and TF); to relate specific variants in transferrin-related genes to brain structure, we determined the additive effect of all variants within these two genes on brain structures that had shown genetic influences in common with transferrin.
Some brain regions were smaller and some larger in people with lower transferrin levels; in fact, there was statistically significant evidence in favor of both effects occurring in different parts of the brain. This pattern of anomalies was somewhat surprising: we expected smaller volumes, not larger ones, in people with low iron (and elevated transferrin) levels. Opposing this, some brain regions involved in neurodegeneration did show lower volumes in those with high iron and low transferrin levels, so iron overload may promote neuronal atrophy in iron-containing structures. The direct association of the volume of these regions to transferrin levels may therefore indicate a future susceptibility to the effects of iron overload and altered transport in these disorders.

Our DTI-based analyses supported a model wherein signs of a less mature or well myelinated brain were found in those with high transferrin levels during adolescence, which is often the liver’s reaction to sustained periods of lower iron availability. The analysis of brain volumes with TBM gives a more complex picture: in segregated comparisons, there were some brain regions that were larger, and some were smaller in those with high transferrin. This imbalance of structure volumes is similar to that seen in some neurogenetic disorders, in which patterns of abnormally high and low volumes are seen (49, 50). As this was not hypothesized, future independent studies are needed to confirm the localization and direction of these effects.

As indicated by a dashed line in Fig. 7, a way to study the iron pathway’s association to brain structure is to determine whether genes influencing transferrin levels also modulate structural variation. Our cross-twin cross-trait genetic analysis revealed that common additive genetic factors influence transferrin concentrations and white matter fiber integrity. Finding neuroanatomical regions whose underlying structure is partially under the same genetic control as transferrin levels can help shed light on the inherited properties of these regions as they develop. Discovering specific iron-associated genetic variants that influence the underlying microstructure in these brain regions could potentially help uncover the neural mechanisms affected by iron transport in the brain. These may lead to downstream genetically mediated impairments.

Specific variants associated with iron mediating proteins in healthy young adults have also been discovered; 40% of the genetic variance in serum transferrin levels is explainable by just a few genetic variants in the TF gene (rs3811647, rs1799852, and rs2280673) and the C282Y mutation in the HFE gene (35). Additionally, interaction between variants in these two genes has been linked to an increased risk of Alzheimer’s disease (51), so the TF and HFE genes are neurobiologically linked. To comprehensively explore these two genes further and determine any coexisting associations to brain structure, we examined all available variants within these genes in regions where the shared additive genetic component between the two traits, transferrin levels, and brain microstructure was statistically significant. We found the H63D mutation within the HFE gene is significantly associated with the mean FA of the left external capsule, one of the regions shown to have significant cross-twin cross-trait correlations. The FA of the external capsule has also been shown to be highly heritable (~60%), however, a sex-by-heritability analysis also shows this region is much more heritable in male subjects (28). Intriguingly, genetic factors also explain a higher proportion of the variance for transferrin levels in men than in women (17).

In a recent study of HFE and TF mutations on iron levels and risk for AD, Giambattistelli et al. (52) found that patients with AD with the H63D polymorphism had increased plasma iron and transferrin levels, but this pattern was not found in healthy control subjects with the variant; in fact, a meta-analysis found that the H63D polymorphism may be protective against AD (53).

As shown in Table S1, the minor allele at rs1799945 (H63D) showed a positive effect on FA. This is the expected direction of association; as mentioned previously, ID can cause deficits in myelin formation, so it is reasonable that an iron overload allele may play a protective role for myelination during neuronal development of these healthy controls.

Our work here is one of the largest bimodal neuroimaging genetics studies of healthy humans to date. It describes a three-step top-down method to analyze gene effects on the brain. First, we related a heritable serum measure—with known cognitive associations—to specific locations in the brain; second, we used a genetic correlation model to hone in on brain regions with evidence of joint genetic determination; and finally, we searched these neuroanatomical locations for variants within genes known to associate with the highly heritable phenotype, serum transferrin. We found localized regions of transferrin association on both the macro and micro scale in the brain. These regions showed additional associations with HFE variants, in a direction that is consistent with several previous studies of brain iron and associated proteins. Future analyses may include studying the mechanistic process of transferrin and HFE in children with cognitive impairment and in elderly subjects with neurodegenerative diseases.

Several conclusions can be drawn about genetic variations that affect transferrin levels and their effect on brain microstructure. Transferrin levels are influenced by at least two known factors: a shortage of iron, which drives them up, and polymorphisms in the two major transferrin-related genes, HFE and TF, as shown in Fig. 7. According to a principle known as Mendelian randomization (54), one can examine genes that are known to affect a measure, such as transferrin, to get a sense of the downstream biological effects of other factors that affect transferrin, such as a shortage of dietary iron. We did find that the H63D mutation within the HFE gene was related to brain FA, but we did not find an effect on FA of the other common (i.e., MAF > 0.05) SNPs in the HFE or TF genes, which are known to explain 40% of variation in transferrin levels. A larger sample may be needed to uncover effects of these SNPs, but a more skeptical alternative interpretation may be that some variants known to powerfully affect transferrin may not affect DTI measures at all. At this point, we cannot say conclusively whether polymorphisms in the TF gene have any causal role on the white matter, although the H63D mutation within the other major transferrin-related gene—HFE—was related to brain FA. A qualified interpretation of the available data would suggest that transferrin levels do relate to brain structure, but further work is needed to clarify which of the several known transferrin-related SNPs, other than HFE H63D, are contributing to the effect.

Methods

Subject Information. A total of 615 subjects (mean age ± SD, 23.5 ± 2.1; 375 women) were included in this study; all subjects had standard structural T1-weighted brain MRI scans and serum iron and transferrin levels measured; 574 also underwent DTI. As part of a reliability analysis, 37 subjects had a duplicate MRI scan taken 3 mo later, and five had a second DTI scan. All subjects were of European ancestry from 350 families. Subjects were recruited as part of a 5-y research project examining healthy young adult Australian twins using structural and functional MRI and DTI with a projected sample size of approximately 1,150 at completion (55). Subjects were screened to exclude cases of pathology known to affect brain structure. No subjects reported a history of significant head injury, a neurological or psychiatric illness, substance abuse or dependence, or had a first-degree relative with a psychiatric disorder. All participants were right-handed as determined using 12 items from the Annett handedness questionnaire (56). We selected only the paired MZ (T1, n = 107; DTI, n = 95 pairs) and same-sex DZ (T1, n = 65; DTI, n = 59 pairs) twins for the cross-twin cross-trait genetic analysis. The rest of the subjects included 52 (n = 43 DTI) pairs of mixed-sex DZ twins, two sets of fraternal triplets (n = 4 individuals), and 112 (n = 95 pairs) individuals unrelated to anyone else in the study; additional subjects included non-twin siblings or unpaired twins with siblings also in the study in which kinship existed between members. A total of 544 of the subjects with

Jahanshad et al.
standard MRI scans were genotyped, of whom 493 also had DTI scans available. Study participants gave informed consent; the studies were approved by the institutional ethics committees in at the University of Queensland and the University of California, Los Angeles. All images used in this analysis went through, and passed, a rigorous quality control; subjects with anatomical abnormalities, imaging artifacts, and misregistered images were removed from analysis and not included in the subject counts.

Blood was collected from subjects at ages 12, 14, and 16 y. Serum was separated from blood samples and stored at −70 °C until assayed; iron, transferrin, and ferritin were measured by using standard clinical chemistry methods (Roche Diagnostics) on a 917 or Modular P analyzer. Data on serum iron and transferrin levels were extracted from these time points and averaged for use in this analysis.

Establishing Zygosity, Genotyping, and Imputation. Zygosity was established objectively by typing nine independent DNA microsatellite polymorphisms (polymorphism information content > 0.7), by using standard PCR methods and genotyping. Results were cross-checked with blood group (ABO, MNS, and Rh) and phenotypic data (hair, skin, and eye color), giving an overall probability of correct zygosity assignment greater than 99.99%. Genomic DNA samples were analyzed on the Human10-Quad BeadChip (Illumina) according to the manufacturer’s protocols (Infinium HD Assay; Super Protocol Guide; Revision A, May 2008). Imputation was performed by mapping the genotyped information to HapMap (release 22, build 36) with Mach software (http://www.sph.umich.edu/csg/abecasis/MACH/index.html).

Image Acquisition. Structural and diffusion-weighted (DW) whole-brain MRI scans were acquired for each subject (4T Medspec; Bruker). T1-weighted images were acquired with an inversion recovery rapid gradient-echo sequence (inversion/repetition/echo times, 700/1500/3.35 ms; flip angle, 8°; slice thickness, 0.9 mm; 256² acquisition matrix). DW images were acquired using single-shot echo-planar imaging with a twice-refocused spin echo sequence to reduce eddy current-induced distortions. A 3-min, 30-gradient acquisition was designed to optimize signal-to-noise ratio for diffusion tensor estimation (57). Imaging parameters were repetition/echo times of 6000/917 ms; field of view, 23 cm; and 128 × 128 acquisition matrix. Each 3D volume consisted of 21 axial slices 5 mm thick with a 0.5-mm gap and 1.8 × 1.8 mm² in-plane resolution. Thirty images were acquired per subject; three with no diffusion sensitization (i.e., T2-weighted b0 images) and 27 DW images (b = 1,132 s/mm²) with gradient directions uniformly distributed on the hemisphere.

Image Preprocessing. Nonbrain regions were automatically removed from each T1-weighted MR image and from a T2-weighted image from the DW image set using FSL software brain extraction tool (58) to enhance coregistration between subjects. All T1-weighted images were corrected for field nonuniformities using FreeSurfer (http://surfer.nmr.mgh.harvard.edu), linearly aligned (with 9 df) to a common space (59). The raw DW images were corrected for eddy current distortions by using the FSL tool ‘eddy_correct’ (http://fsl.fmrib.ox.ac.uk/fsl/). For each subject, the three eddy-corrected images with no diffusion sensitization were averaged, linearly aligned, and resampled to the subject’s corresponding down-sampled T1 image. The average b0 images were then elastically registered to the subject’s aligned T1-weighted structural scan by using an inverse consistent registration with a mutual information cost function (60) to adjust for any echo-planar-induced susceptibility artifacts.

TBM. TBM is a technique that identifies regional structural differences from the gradients of the deformation fields that align brain images to a common anatomical template. After nonlinearly aligning the full brain of all subjects to their corresponding minimum deformation template (MDT), a separate Jacobian map (i.e., relative volume map) was created for each subject. These Jacobian maps share the coordinate system defined by the MDT, help to characterize the local volume differences between one individual and the normal anatomical template. These maps explain the relative expansion and contraction of regions from each individual relative to the template.

Computing Anisotropy and Diffusivity. Under a single-tensor model (61), diffusion of water molecules attenuates the MR signal in direction r, according to the Stejskal–Tanner equation:

\[
S_r(t) = S_0(e^{-tD_r(t)})
\]

Here, \(S_r(t)\) is the non-DW baseline intensity in direction r, \(D_r(t)\) is the apparent diffusion coefficient, and \(S_0\) is a constant depending on the gradient k. Diffusion tensors were computed from the 27-gradient DW images using FSL software (http://fsl.fmrib.ox.ac.uk/fsl/). The FA of diffusion was computed from the tensor eigenvalues \(\lambda_1, \lambda_2, \lambda_3\) at each voxel. FA is influenced by both axial diffusivity \(D_A\) (a measure of diffusion along the axonal fibers) and radial diffusivity (the average of \(\lambda_2\) and \(\lambda_3\), a measure of diffusion orthogonal to the axonal fibers).

Template Creation and Registration. We created a MDT by using nonlinear fluid registration (62), with the method proposed by Kochunov and colleagues (63, 64). The N 3D vector fields fluidly registering a specific individual to all other N participants were averaged and applied to that subject. This geometrically adjusts the anatomy but preserves the intensities and anatomical features of the template subject.

To create a representative MDT for the TBM analysis, we randomly selected 32 (16 female/16 male) nonrelated participants’ T1-weighted images, after dosing-correcting and brain morpbometry. The images were then registered to the Colin template (59), and created two MDTs with 16 subjects (eight female) in each group. For one group, the target was male and the other female. The templates for each group were then averaged to create one representative anatomically centered target. Skull-stripped T1-weighted images for each subject were registered to the final population averaged FA-based MDT by using an inverse consistent 3D elastic warping technique using a mutual information cost function (60).

To create an MDT for DTI analysis, we selected the same 32 participants, yet we used their FA images (calculated after \(b_0\) susceptibility correction) to create the MDT in the exact same manner as the T1-weighted structural scans. FA maps for each of the susceptibility corrected subjects were registered to the final population averaged FA-based MDT by using a 3D elastic warping technique with a mutual information cost function (60). To further align white matter regions of interest, the FA-based MDT and all whole-brain registered FA maps were then thresholded at 0.25, as FA values lower than 0.25 in healthy-appearing white matter may reflect contributions from nonwhite matter. Individual thresholded FA maps were then reregistered to the thresholded MDT in the same way as the whole brain registration. After filtering of the FA maps, the FA images were smoothed with a Gaussian filter with an isotropic full-width half maximum of 5 mm.

Random-Effects Regression. The relationships of transferrin to measures of iron and transferrin brain morphometry were assessed at each voxel in the brain by using a mixed-effects regression model to account for similarities within families while controlling for the effects of sex, age, and serum iron levels. To boost power of the association, and reduce random noise brought on by image acquisition, we included duplicate scans for the subjects who had them available. The variable of interest (transferrin), sex, and age were included as fixed effects. Random intercepts were included for each family and subjects to account for relatedness within families as well as the duplicate scans used. The analysis was implemented in the R statistical package (version 2.9.2; http://www.r-project.org/) using the ‘lme’ library (65).

Two-tailed hypotheses. As noted earlier, extremely high and extremely low \(b_0\) levels of iron can adversely affect the brain, but these observations do not completely imply in which direction the correlation would be in healthy people who maintain their iron levels for the most part in the normal range. As iron overload was not expected in this young, healthy population, mild insufficiencies in iron were considered more likely. This led to a directional hypothesis that poorer brain phenotypes might be found in those with lower chronic levels of iron (as inferred from transferrin measures). However, we considered it also possible, but less plausible, that there might be enough people with very high iron levels to drive the effect in the opposite direction. To allow for this alternative but less likely hypothesis, we ran our analyses with a more conservative searchlight FDR threshold of 0.025, to allow us to reject the null hypothesis in either direction, but distinguish between the alternative hypotheses in different directions. More information may be found in SI Methods.

Cross-Twin Cross-Trait Analysis. We used a cross-twin cross-trait analysis (66) to detect common genetic or environmental factors influencing both brain structure (or microstructure) and serum transferrin levels at every voxel within the brain. Covariance matrices for the phenotypes, in this case the voxel-wise structural measure of interest (structural deformation, microstructural anisotropy, or diffusivity) and serum transferrin levels were computed between the MZ twins who share all the same genes, and the DZ twins who share, on average, half of their genetic polymorphisms. These covariance matrices were then entered into a multivariate structural equation modeling (SEM) (67), using OpenMX software (http://openmx.psych.virginia.edu) to fit the relative contributions of additive genetic (A), shared environmental (C), and unshared or unique environmental (E) components to
the population variances and covariances of the observed variables. Experimental mice were included in the E component, as assumed to be independent between twins 1 and 2 (i.e., no correlation).

In multivariate SEM, it is assumed that there are common genetic and environmental factors that affect various phenotypes. Here we consider bivariate models with two phenotypes, transferrin levels and the brain MRI- or DTI-derived value at each voxel. The common genetic and environmental components of the variance may be estimated from the total population variance by examining the difference between the covariances of the MZ and DZ twins within the same individual (cross-trait within individual) and also between one phenotype in one twin with the other phenotype in the second twin (cross-trait cross-trait). By using this multivariate SEM, we can also obtain the additive genetic and shared environmental influences on the correlations between the two phenotypes, denoted as \(r_a\) and \(r_c\), respectively. A path diagram describing the SEM and the connections between the twins is shown in Fig. 8.

The cross-trait within-individual correlation [i.e., the correlation between the voxel value (V) and transferrin (T) in twin 1 or in twin 2] is divided into additive genetic and shared and unique environmental components (e.g., \(A_x\), \(C_x\), and \(E_x\) for voxel value and \(A_y\), \(C_y\), and \(E_y\) for transferrin; \(i = 1\) or 2), and the correlation coefficients between \(A_{ij}\) and \(A_{ij}\), \(C_{ij}\) and \(C_{ij}\), and \(E_{ij}\) and \(E_{ij}\) are denoted by \(r_{a_{ij}}\), \(r_{c_{ij}}\), and \(r_{e_{ij}}\) respectively. The cross-trait cross-trait correlation is shown as \(A_{ij}\) and \(A_{ij}\), and \(C_{ij}\) and \(C_{ij}\) for the voxel value in twin 1 and the transferrin level in twin 1, where \(i = 1\) or 2 and \(j \neq i\).

There is no \(r_{a_{ij}}\) term for \(E_x\) and \(E_y\), because the unique factors between subjects are independent. The covariance across the two phenotypes within the same subject, or separately in the two subjects, is then derived by multiplication of the path coefficients for the closed paths in the path diagram. For example, covariance between the voxel values in twin 1 and the transferrin level in twin 2 is equal to \(a_r v^{(1)} a_r + c_r c_r + e_r e_r\) for MZ twins, and \(a_r v^{(1)} a_r + c_r c_r + e_r e_r\) for DZ twins. This implies that any excess in cross-trait cross-trait correlation in MZ twins over that in DZ twins is attributed to common genetic factors that affect both voxel values and transferrin levels.

Paths drawn between the same phenotype would be identical to considering a univariate voxel-wise SEM model (27). For \(A_1\) and \(A_2\), the correlation coefficient is equal to 1 for MZ and 0.5 for DZ twin pairs. The correlation coefficient between \(C_1\) and \(C_2\) is always 1 from the definition of the shared environment, and \(E_1\) and \(E_2\) are assumed to be independent and there is no correlation.

In twin studies, it is common to examine whether the observed measures are best modeled by using a combination of additive genetic and shared and unshared environmental factors, or whether only one or two of these factors is sufficient to explain the observed pattern of inter-twin correlations.

If the correlation between the voxel value of the image in one twin and the level of transferrin in the other twin is greater in MZ pairs than in DZ pairs, then, under standard assumptions, the greater correlation may be assumed to be caused by common genetic factors controlling both factors. In the univariate model with a single phenotype, which we denote \(x\), the genetic and environmental contributions in twin \(j = 1\) or 2) is modeled by defining the following:

\[
\begin{align*}
  z_j &= a_r x_j + c_r c_j + e_r e_j \\
  a_r, c_r, \text{ and } e_r &\text{ denote the additive genetic and shared and unshared environmental components. Cross-trait correlations between voxel values (v) and serum transferrin (t) level are then derived from the covariance matrix of the following vector: }
  \begin{pmatrix}
    \mathbf{V}_1 \\
    \mathbf{T}_1 \\
    \mathbf{V}_2 \\
    \mathbf{T}_2
  \end{pmatrix}
\end{align*}
\]

given by the following 4 \(\times\) 4 matrix:

\[
\begin{align*}
  \Phi_v &= \begin{pmatrix}
    \text{cov}(v_1, t_1) & \text{cov}(v_1, t_2) \\
    \text{cov}(v_2, t_1) & \text{cov}(v_2, t_2)
  \end{pmatrix} \\
  \Phi_v &= \begin{pmatrix}
    \text{cov}(v_1, t_1) & \text{cov}(v_2, t_2) \\
    \text{cov}(v_2, t_1) & \text{cov}(v_1, t_2)
  \end{pmatrix}
\end{align*}
\]

where \(\Phi_v\) and \(\Phi_t\) are the 2 \(\times\) 2 covariance matrices for phenotype \(v\) or \(t\) between twins 1 and 2, as performed in univariate SEM. \(\Phi_v\) is the cross-trait covariance matrix, composed of the covariance between the two traits within the different unrelated individuals [\(\text{cov}(v_1, t_1)\) and \(\text{cov}(v_2, t_2)\)] and the cross-trait cross-trait covariance between the pairs [\(\text{cov}(v_1, t_2)\) and \(\text{cov}(v_2, t_1)\)], as detailed below:

\[
\Phi_v = \begin{pmatrix}
  \text{cov}(v_1, t_1) & \text{cov}(v_1, t_2) \\
  \text{cov}(v_2, t_1) & \text{cov}(v_2, t_2)
\end{pmatrix}
\]

where \(\alpha_1\) is 1 for MZ twins, and 0.5 for DZ twins, \(r_a\) and \(r_c\) are the cross-trait correlation coefficients for \(A\) and \(C\) and \(E\) for \(v\) and \(t\), respectively. A higher value of \(r_v\) indicates that the two phenotypes are more likely mediated by a common set of genes (34, 68). The path coefficients were estimated by comparing the covariance matrix implied by the model and the sample covariance matrix of the observed variables, using maximum-likelihood fitting to give a \(\chi^2\) value. We started from the full set of path coefficients \((a_r, c_r, e_r, a_t, c_t, E_v, E_r)\) and removed one of \(a_r\), \(c_r\), \(e_r\), or \(a_t\), \(c_t\) or \(E_r\) or \(E_v\) or \(E_t\) at a time. The likelihood-ratio test was then performed. If the likelihood ratio is positive, \(a_r\) or \(c_r\) or \(e_r\) is removed from the model, and otherwise \(a_t\), \(c_t\), or \(E_v\), or \(E_t\) or \(E_r\) is removed from the model, and otherwise the model remains the same.

If two models contained the same number of parameters, the model with a smaller \(\chi^2\) value was considered better. Model selection ended when the best model was achieved, i.e., when either (i) all possible more restricted models were not better than the current model or (ii) the current model was the most restricted and contained \(e_r\), \(e_v\), and \(r_v\) only. If \(r_v\) was included in the best model, the significance of \(r_v\) was then determined by comparing the \(\chi^2\) values of the best model and its submodel where \(r_v\) is 0. To determine the significance of the submodels, or the restricted models, with respect to the full model, we obtain the log-likelihood for the full and the restricted models, denoted by \(\log \text{L}(\text{full})\) and \(\log \text{L}(\text{sub})\), respectively. Minus two times this difference, or \(-2(\log \text{L}(\text{full}) - \log \text{L}(\text{sub}))\), is asymptotically distributed as a \(\chi^2\) distribution with the df equal to the difference between the df of the two models, and therefore the inverse \(\chi^2\) distribution is estimated with these parameters.

SNP Selection. We examined all SNPs within two genes previously shown (35) to affect serum transferrin levels in healthy adults: TF and HFE. By using HapMap, we searched for SNPs that met our criteria of having an MAF greater than 5%; when matching these SNPs to those with available genotype information in our imputed data, 42 valid SNPs from these genes were available for analysis. We determined significance levels for association tests by first examining the total number of independent tests performed. Linkage disequilibrium among SNPs tested corresponds to correlation between the SNPs, and therefore each test is not completely independent. By first estimating the effective number of independent tests, we can avoid using a significance level too conservative for the number of tests we performed. As a result of linkage disequilibrium, the effective number of SNPs tested (38, 39) was 20.

ROI SNP Association in Significantly Correlated Clusters. In each cluster (\(\geq 27\) voxels to represent a size equivalent to one voxel and all its surrounding neighbors) that was found to have significant cross-trait cross-twin additive genetic associations, we found the average value across all of the voxels in that region and performed univariate associations with all 42 SNPs by using a mixed-model approach controlling for age, sex, and serum iron levels.
Multiple Comparisons Correction. Computing thousands of tests of association on a voxel-wise level can introduce a high type I (i.e., false-positive) error rate in neuroimaging studies. To control these errors, we used a searchlight method for FDR correction as described by Langers et al. (70), which ensures a regional control over the FDR in any reported findings. To ensure adequate regionally selective associations with the transferrin levels, we use the standard FDR (37, 71) procedure as opposed to searchlight FDR, as we would like to determine the best overall fit of the SEM model, and not necessarily examine any localized or clustering effects. When examining the significance of the effects of the SNPs regressed on the mean FA value within ROIs with high correlation, coexistent associations, we corrected for multiple comparisons by using the strict Bonferroni correction controlled at the q level of 0.05, at which a threshold for significance was determined by dividing 0.05 by the effective number of SNPs tested (20), and the number of ROIs these SNPs were each tested previously (5). The Bonferroni threshold for significance was therefore set as follows:

\[ q = 0.05 / (20 \times 6) = 0.00042 \]

Post Hoc Analysis: Voxel-Wise Effect of HFE H63D Polymorphism on Fiber Integrity. The number of minor alleles for each HFE H63D (rs1799945) was regressed against the FA at each voxel within the white matter, after adjusting for sex and age as before. Family structure was taken into account using mixed-effects modeling (72). To correct for multiple comparisons across voxels, we used a searchlight method to control the FDR regionally (70).

ACKNOWLEDGMENTS. We thank the twins and siblings for their participation. In Brisbane, we thank Marlene Grace and Ann Eldridge for twin recruitment, Aiman Al Najjar and other radiographers for scanning, Kori Johnson for scanning and data transfer, Kerrie McAloney and Daniel Park for research support, and staff in the Molecular Epidemiology Laboratory for serum and DNA sample processing and preparation. This work was supported by National Institute of Child Health and Human Development Grant R01 HD050735, National Health and Medical Research Council (NHMRC; Australia) Grant 486682, Grant T15 LM07336 (to N.J.), The Achievement Rewards for College Scientists Foundation (J.L.S.), National Institute of Mental Health Grant 1F31MH078061 (to J.L.S.), and Australian Research Council Future Fellowship FT0991634 (to G.J.d.Z.). Genotyping was supported by NHMRC Grant 389875. Additional support for algorithm development was provided by National Institutes of Health Grants R01 EB004832, R01 EB008281, and R01 EB007813.


