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 Genetic Contributions to Human Brain Morphology and Intelligence

Hilleke E. Hulshoff Pol,1 Hugo G. Schnack,1 Danielle Posthuma,2 René C. W. Mandl,1 Wim F. Baaré,1 Clarine van Oel,1 Neeltje J. van Haren,1 D. Louis Collins,3 Alan C. Evans,1 Katrin Amunts,4,5 Uli Bürgel,6,7 Karl Zilles,6,7 Eco de Geus,2 Dorret I. Boomsma,2 and René S. Kahn1

1Department of Psychiatry, Rudolf Magnus Institute of Neuroscience, University Medical Center Utrecht, The Netherlands, 2Department of Biological Psychology, Free University Amsterdam, 1081 BT Amsterdam, The Netherlands, 3Montreal Neurological Institute, McGill University, Montreal, Quebec, Canada, H3A 2B4, 4Institut fuer Medizin, Forschungszentrum Juelich, D-52405 Juelich, Germany, Departments of 5Psychotherapy and Psychiatry and 6Neurosurgery, Die Leitseite der Rheinisch-Westfa¨lischen Technischen Hochschule Aachen University, D-52062 Aachen, Germany, and 7C. and O. Vogt Institute for Brain Research, Heine University Du¨sseldorf, D-40225 Du¨sseldorf, Germany

Variation in gray matter (GM) and white matter (WM) volume of the adult human brain is primarily genetically determined. Moreover, total brain volume is positively correlated with general intelligence, and both share a common genetic origin. However, although genetic effects on morphology of specific GM areas in the brain have been studied, the heritability of focal WM is unknown. Similarly, it is unresolved whether there is a common genetic origin of focal GM and WM structures with intelligence. We explored the genetic influence on focal GM and WM densities in magnetic resonance brain images of 54 monozygotic and 58 dizygotic twin pairs and 34 of their siblings. For genetic analyses, we used structural equation modeling and voxel-based morphometry. To explore the common genetic origin of focal GM and WM areas with intelligence, we obtained cross-trait/cross-twin correlations in which the focal GM and WM densities of each twin are correlated with the psychometric intelligence quotient of his/her cotwin. Genes influenced individual differences in left and right superior occipitofrontal fascicle (heritability up to 0.79 and 0.77), corpus callosum (0.82, 0.80), optic radiation (0.69, 0.79), corticospinal tract (0.78, 0.79), medial frontal cortex (0.78, 0.83), superior frontal cortex (0.76, 0.80), superior temporal cortex (0.80, 0.77), left occipital cortex (0.85), left postcentral cortex (0.83), left posterior cingulate cortex (0.83), right parahippocampal cortex (0.69), and amygdala (0.80, 0.55). Intelligence shared a common genetic origin with superior occipitofrontal, callosal, and left optical radiation WM and frontal, occipital, and parahippocampal GM (phenotypic correlations up to 0.35). These findings point to a neural network that shares a common genetic origin with human intelligence.

Key words: genetics; twins; neuroanatomy; voxel-based morphometry; intelligence; cognition

Introduction

Variation in total gray and white matter volume of the adult human brain is primarily (70–80%) genetically determined (Pennington et al., 2000; Pfeifferbaum et al., 2000; Baaré et al., 2001). Moreover, total brain volume is positively correlated with general intelligence, and both share a common genetic origin (Posthuma et al., 2002). However, this finding does not necessarily mean that genes influence focal brain structures in the same manner throughout the brain. Moreover, it does not necessarily mean that a common genetic origin with general intelligence is shared by all structures throughout the brain.

Determining the extent to which focal brain morphology and its association with cognitive functioning are influenced by genes (or environment) is important for both our understanding of healthy functioning and elucidating the causes of brain disease. More specifically, it enhances our knowledge of individual variation in brain functioning and aids in the interpretation of the morphological changes as found in psychiatric disorders such as schizophrenia. Also, it allows future efforts to find particular genes responsible for brain structures and functions to be concentrated on brain areas that are under considerable genetic influence.

To date, most studies focused on the genetic influence on variation in total brain volume. A few studies have examined genetic effects on more specific brain areas throughout the brain. Morphology of focal frontal and temporal gray matter (GM) areas is particularly influenced by genetic factors (Thompson et al., 2001; Wright et al., 2002). However, the genetic influence on focal white matter (WM) has not been examined. It has been suggested that frontal GM volume and intelligence share genetic factors, but it also remains unresolved whether this association...
shares a common genetic origin (Thompson et al., 2001; Toga and Thompson, 2005).

To determine the relative contributions of genetic, common, and unique environmental influences on variation in focal brain structures and their common origin with intelligence, the (extended) twin model is a powerful approach (Posthuma and Boomsma, 2000), particularly because morphological (Hulshoff Pol et al., 2002) and cognitive (Posthuma et al., 2000) findings in twins can be extended toward the general singleton population. For genetic influences, the extent to which monozygotic (MZ) twin pairs resemble each other more than is the case for dizygotic (DZ) twin pairs is the determining factor. However, in addition to genetic influences, common (or shared) environmental influences may play a role in explaining resemblances. The presence of shared environmental factors is suggested when correlations in DZ twins are larger than half the MZ correlation (Boomsma et al., 2002). A first impression of the importance of unique environmental factors is obtained from the extent to which MZ twins do not resemble each other. In a similar manner, the extent to which genetic and environmental factors influence the association of two traits such as brain structure and intelligence can be determined by comparing the cross-trait/cross-twin correlation between brain structure and intelligence in MZ and DZ twins. If cross-trait/cross-twin correlations are larger in MZ twins, this suggests genetic mediation of the association.

Materials and Methods
Magnetic resonance brain images were analyzed using voxel-based morphometry (VBM) (Collins et al., 1995; Hulshoff Pol et al., 2001, 2004). The extent to which GM and WM densities of MZ twin pairs correlate with each other compared with the correlations of DZ twin pairs and siblings enables the calculation of brain maps of the genetic contribution to focal GM and WM areas throughout the brain. For the genetic analyses, structural equation modeling (SEM) was used implemented in Mx software (Neale, 1997) in each voxel separately. The significance level was corrected for multiple comparisons according to random field theory (Worsley et al., 1996; Ashburner and Friston, 2000). To define which anatomical GM areas were involved, the genetic WM maps were superimposed onto histologically defined human anatomical WM brain maps (Bürgel et al., 1999; Rademacher et al., 2001). The cross-trait/cross-twin correlations were obtained between focal GM and WM density and intelligence, verbal (VIQ) and nonverbal (performance) (PIQ) intelligence quotient measures of the Dutch version of the Wechsler Adult Intelligence Scale III-Revised (WAIS-IIIR). We chose to include VIQ and PIQ separately because they correlate.

When interpreting the findings, limitations of our chosen approach have to be considered. The VBM procedure involves several steps. VBM requires registration of the separate brain images onto a model brain to allow for statistical inferences in focal brain areas. The choice of the model brain may have influenced the results. However, when comparing several model brains with a similar dataset in a previous study, we found that the results remained the same. Thus, the influence of the model brain on our results seems to have been of minimal influence. Moreover, registration error will occur during this procedure and is unavoidable. Therefore, some caution in interpreting these results is warranted. However, because we registered each individual brain separately to the model brain, it seems unlikely that a bias occurred that behaved differently in MZ and DZ twins other than the similarities that we expected to find attributable to differences in the overlap of their genetic makeup. Also, note that, because we were especially interested in the influence of genes on focal brain densities, thus after correction for the known overall effect of genes on GM and WM volumes, we did not include a volume-preserving step (Good et al., 2001a) in the procedure. Thus, the GM and WM that we now find to be highly heritable include those areas in which the individual variation is influenced by genes on top of the genetic influences on global brain volume. One also has to keep in mind that the focal brain areas that we do not find to be genetically determined may still be determined by genes that influence overall brain volume. Alternatively, these focal brain areas may be influenced by environmental factors. Finally, our overall age correction may be only relevant for the age range as included in the sample. Aged and early development samples can and probably will give different patterns. Recently, a longitudinal study in children and adolescent singletons revealed a nonlinear trajectory of change in cortical thickness, which was associated with intelligence (Shaw et al., 2006). Within the age range of 19–69 years, changes in the genetic and environmental influences on brain morphology and in the relationship between brain morphology and intelligence may also occur. Analyses of possible changes in genetic and environmental influences with age await longitudinal follow-up measurements in these subjects.

Demographics.
For magnetic resonance brain imaging, a total number of 258 family members from 112 families participated in the study after written consent was obtained (Baare´ et al., 2001) (Table 1). They consisted of 33 MZ male, 17 DZ male, 21 MZ female, 20 DZ female, and 21 DZ opposite-sex twin pairs, and 19 male and 15 female full siblings. Subjects were between 19 and 69 years of age (mean ± SD, 30.7 ± 9.6 years). Handedness was determined by the Cognitive Assessment for Handedness and Laterality Data from the Comprehensive Assessment of Symptoms and History (Andreasen et al., 1992): 213 subjects were right-handed, 34 were left-handed, and 11 were mixed-handed. Twins were recruited from the (healthy) twin sample of the Department of Psychiatry of the University Medical Centre Utrecht and the Netherlands Twin Registry (http://www.tweelingenregister.org/index_uk.html). DNA testing using the polymorphic markers D06S474, D07S1804, D07S1870, D12S811, D13S119, D13S126, D13S788, D20S119, D22S683, DXS1001, and ELN, or D13S317, VWA, D7S452, D35158, TH01, TP0X, CSF1P0, and D5S818 determined zygosity. Except for one twin pair, all twins and their siblings were reared together. Two twin pairs were born by cesarean section delivery. Mental and physical health was assessed by means of the Family Interview for Genetic Studies (Nurnberger et al., 1994) and a medical history inventory, respectively. All subjects’ consent was obtained according to the declaration of Helsinki. The Scientific and Ethical Committee of the University Medical Centre Utrecht, in which the study was performed, approved the study.

Image acquisition.
Magnetic resonance images were acquired on a Phillips (Amsterdam, The Netherlands) NT scanner operating at 1.5 T in all subjects. T1-weighted three-dimensional (3D) fast-field echo scans with 160–180 1.2 mm contiguous coronal slices (echo time (TE), 4.6 ms; repetition time (TR), 30 ms; flip angle, 30°; field of view (FOV), 256
Table 1. Demographics

<table>
<thead>
<tr>
<th></th>
<th>MZ</th>
<th>DZ</th>
<th>SIB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Individuals (MRI/IQ)</td>
<td>49/213</td>
<td>65/285</td>
<td>34/190</td>
</tr>
<tr>
<td>Sex, M/F (MRI)</td>
<td>34/15</td>
<td>33/32</td>
<td>19/15</td>
</tr>
<tr>
<td>Sex, M/F (IQ)</td>
<td>98/115</td>
<td>117/168</td>
<td>91/99</td>
</tr>
<tr>
<td>Hand, R/L, A (MRI)</td>
<td>41/8</td>
<td>54/11</td>
<td>29/5</td>
</tr>
<tr>
<td>Hand, R/L, A (IQ)</td>
<td>183/30</td>
<td>251/34</td>
<td>175/15</td>
</tr>
<tr>
<td>Age, years (MRI)</td>
<td>28.3 (3.6)</td>
<td>29.2 (8.7)</td>
<td>29.6 (4.8)</td>
</tr>
<tr>
<td>Hand, years (IQ)</td>
<td>37.7 (36.6)</td>
<td>36.6 (13.3)</td>
<td>36.7 (13.2)</td>
</tr>
<tr>
<td>IQ (IQ)</td>
<td>63.3 (13.3)</td>
<td>64.9 (12.2)</td>
<td>65.4 (13.3)</td>
</tr>
<tr>
<td>IQ (IQ)</td>
<td>76.0 (12.8)</td>
<td>75.0 (14.0)</td>
<td>75.6 (13.2)</td>
</tr>
</tbody>
</table>

Values are shown as mean (SD). M, Male; F, female; R, right; L, left; A, ambidexter; SIB, singleton siblings of twin pairs.
anterior (positive values). The in the left and right medial frontal cortex and right superior frontal cortex. A heritability with a peak value of 0.83 was found for GM density in the left occipital cortex, and of 0.83 in the left posterior cingulate (Table 2, Fig. 1) (see Fig. 4). In addition, heritabilities of 0.80 and 0.55 were found in the left and right amygdala, respectively, and of 0.69 in the right parahippocampal gyrus. Heritabilities with a peak value of 0.79 were found for WM density in the superior occipitofrontal fascicle bilaterally, of up to 0.82 throughout the corpus callosum in the left and right hemisphere.

**Results**

**Sources of variation in focal GM and WM areas**

A heritability with a peak value of 0.83 was found for GM density in the left and right medial frontal cortex and right superior frontal cortex, of 0.80 and 0.77 in the left and right Heschl’s gyrus, of 0.83 in the left occipital cortex, and of 0.83 in the left posterior cingulate (Table 2, Fig. 1) (see Fig. 4). In addition, heritabilities of 0.80 and 0.55 were found in the left and right amygdala, respectively, and of 0.69 in the right parahippocampal gyrus. Heritabilities with a peak value of 0.79 were found for WM density in the superior occipitofrontal fascicle bilaterally, of up to 0.82 throughout the corpus callosum in the left and right hemisphere.

**Statistical analysis of common origin between GM and WM density and intelligence.** To test whether verbal (VIQ) and nonverbal (PIQ) intelligence and GM and WM density share a common origin attributable to genetic, common, or unique environmental factors, the “observed correlation” ($r_{O}$) between GM and WM density and VIQ/PIQ was decomposed into genetic and environmental components, but only in those voxels in which density was significantly determined by genetic factors.

Decomposition of the association was based on the comparison of cross-trait/cross-twin correlations for MZ and DZ twins (or sibling pairs) (Posthuma et al., 2002). If the correlation between GM and WM density of twin 1 and VIQ or PIQ of twin 2 is larger in MZ than in DZ twins, this indicates that the genes influencing GM and WM density partly overlap with the genes that influence VIQ/PIQ. The extent of the overlap is reflected by the magnitude of the genetic correlation ($r_{G}$). $r_{G}$ reflects the correlation between the set of genes that influences focal GM and WM density and the set of genes that influences VIQ/PIQ. When the cross-trait/cross-twin correlations are similar for MZ and DZ twins, this suggests that environmental factors contribute to the observed phenotypic correlation between GM and WM density and VIQ/PIQ. Given a heritability of 0.80 for brain density and a heritability of 0.80 for VIQ/PIQ and a correlation between brain volume and VIQ/PIQ of 0.25, at least 16 MZ and 16 DZ twin pairs are needed.

In 24 MZ pairs, 31 DZ pairs, and 25 additional siblings (in these 135 subjects, both volumetric and IQ data were available), the correlations between GM and WM density and VIQ/PIQ were decomposed into genetic (A) and unique environmental components (E) using structural equation modeling for a trivariate genetic design for GM and WM density separately. Observed correlations with absolute values below 0.10 were not considered for additional analysis.

**Table 2. Heritability estimates for focal gray and white matter areas with significant genetic contribution**

<table>
<thead>
<tr>
<th>Region</th>
<th>Heritability</th>
<th>Talairach</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gray matter</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Superior frontal</td>
<td>0.76</td>
<td>-5</td>
<td>58</td>
<td>35</td>
<td></td>
</tr>
<tr>
<td>Medial frontal</td>
<td>0.78</td>
<td>-37</td>
<td>34</td>
<td>39</td>
<td></td>
</tr>
<tr>
<td>Postcentral gyrus</td>
<td>0.83</td>
<td>-55</td>
<td>-29</td>
<td>57</td>
<td></td>
</tr>
<tr>
<td>Posterior cingulate</td>
<td>0.83</td>
<td>-5</td>
<td>-33</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>Heschl’s gyrus</td>
<td>0.80</td>
<td>-27</td>
<td>-31</td>
<td>21</td>
<td></td>
</tr>
<tr>
<td>Amygdala</td>
<td>0.80</td>
<td>-29</td>
<td>-5</td>
<td>-7</td>
<td></td>
</tr>
<tr>
<td>Occipital cortex</td>
<td>0.85</td>
<td>-25</td>
<td>-60</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td><strong>White matter</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Superior occipitofrontal fascicle</td>
<td>0.79</td>
<td>-17</td>
<td>10</td>
<td>25</td>
<td></td>
</tr>
<tr>
<td>Corpus callosum</td>
<td>0.82</td>
<td>-21</td>
<td>-9</td>
<td>35</td>
<td></td>
</tr>
<tr>
<td>Optic radiation</td>
<td>0.69</td>
<td>-35</td>
<td>-48</td>
<td>17</td>
<td></td>
</tr>
<tr>
<td>Corticospinal tract</td>
<td>0.78</td>
<td>-17</td>
<td>-12</td>
<td>21</td>
<td></td>
</tr>
</tbody>
</table>

*The $r$ values indicate the distance in millimeters from the left (negative values) to the right (positive values) side of the brain passing through the anterior commissure. The $y$ values indicate the distance from posterior (negative values) to anterior (positive values). The $z$ values indicate the distance from inferior (negative values) to superior (positive values). Two separate genetically determined areas were identified within the superior and medial frontal cortices.

**Figure 1. Genetically influenced focal GM density brain areas.** Heritability estimates of GM density in focal brain areas in healthy adult humans are shown for the significance level thresholded A-map superimposed on axial and sagittal sections through the magnetic resonance image of the standardized reference brain (left) and for the complete A-map (right). The significance level thresholded A-map shows the voxels that had a significant fit of the genetic model compared with the other models based on the likelihood ratio test and using structural equation modeling. Moreover the level of significance was set at $\chi^2 = 25.3$ for $df = 1$ and $\chi^2 > 29.3$ for $df = 2$ after correction for multiple comparisons according to random field theory (the uncorrected significance levels would have been $\chi^2 > 3.8$ and $\chi^2 > 6.0$). Analyses were controlled for age, sex, and handedness. a, Left occipital cortex. b, Right medial frontal gyrus. c, Left and right Heschl’s gyrus. d, Left amygdala and right parahippocampal gyrus. R, Right.
Itations are $r_p = 0.25$, $r_g = 0.37$, occipital cortex (positively associated with PIQ, $r_p = 0.17$, $r_g = 0.23$), and right parahippocampal gyrus (positively associated with PIQ, $r_p = 0.23$, $r_g = 0.40$; positively associated with VIQ, $r_p = 0.11$, $r_g = 0.19$) (Table 3, Fig. 5). Also, a shared genetic origin was suggested between WM density and VIQ/PIQ in parts of the superior occipitofrontal fascicle with genetically determined associations that were positive for PIQ on the left ($r_p = 0.17$, $r_g = 0.24$) and on the right ($r_p = 0.35$, $r_g = 0.32$). In addition, positive genetically determined associations were found in the corpus callosum on the left with PIQ ($r_p = 0.18$, $r_g = 0.22$) and on the right with VIQ ($r_p = 0.14$, $r_g = 0.15$), as well in the optic radiation on the left with PIQ ($r_p = 0.26$, $r_g = 0.39$). Note that there were positive genetic associations with VIQ in the superior occipitofrontal fascicle bilaterally. However, the strict criterion set in the study prevented these genetically determined associations from reaching statistical significance.

**Discussion**

Our data indicate that, even after correcting for overall brain size (through scaling brains during the transformation procedure), and thus for the considerable genetic influences on total brain volume, and after stringent correction for multiple comparisons of the voxel-based genetic analyses, we found genes to significantly influence WM density of the superior occipitofrontal fascicle, corpus callosum, optic radiation, and corticospinal tract, as well as GM density of the medial frontal, superior frontal, superior temporal, occipital, postcentral, posterior cingulate, and parahippocampal cortices. Moreover, our results show that VIQ/PIQ share a common genetic origin with an anatomical neural network involving the frontal, occipital, and parahippocampal GM and connecting GM of the superior occipitofrontal fascicle, and corpus callosum.

Variation in the superior occipitofrontal fascicle was found to be highly heritable. Voxel-based morphometry analysis on anatomical magnetic resonance images does not allow for white matter tract tracing, such as is possible with diffusion tensor images. However, the majority of the WM voxels that we found to be highly heritable did essentially overlap with and follow the anatomical pathway of the superior occipitofrontal fascicle as shown by its superposition onto the PM histological probability maps. In contrast, relatively few voxels overlapped with the probability maps of other white matter tracts. Moreover, there was no indication that these voxels followed, for example, the right-to-left hemisphere direction of the corpus callosum, the superior-to-inferior direction of the corticospinal tract, or the distinguished pattern of the optic tract.

The anatomical and functional knowledge of the superior occipitofrontal fiber tract, as with many other brain fiber tracts, is still sparse. However, the GM areas that we (and others) found to

**Figure 2.** Genetically influenced focal WM density brain areas superimposed on the histologically defined map of the superior occipitofrontal fascicle. Heritability of WM density in focal brain areas in healthy adult humans is shown for the significance level thresholded A-map superimposed on axial and sagittal sections in the left (L) and right (R) hemisphere through the magnetic resonance image of the standardized reference brain (left) and superimposed on the histologically defined map of the occipitofrontal superior fascicle (middle) (reproduced with the permission of K.Z, K.A., and U.B.). The complete A-maps are shown on the right. Note that several of the genetically influenced focal WM density voxels shown on the left also overlapped with the histologically defined maps of the corpus callosum, optic radiation, and corticospinal tract, which are not shown in the middle figures because relatively few voxels overlapped with these white matter tracts. a. Axial section. b. c. Sagittal sections.
be highly heritable may communicate through this pathway, forming a neural network. The superior occipitofrontal fascicle is a corticocortical association pathway. Fibers arise in the lateral prefrontal cortex of the inferior and middle frontal gyri and join to form a tight bundle at the level of the anterior horn of the lateral ventricle. The tract then runs posterior, superolateral to the caudate and can be easily seen in coronal sections at the apex of the lateral ventricle (Dejerine, 1895). It is considered to carry fibers that connect occipital, temporal (Crosby et al., 1962), and parietal (Cantani et al., 2002) regions with the frontal cortex. In monkeys, this tract is known to project to areas 46 and 8 (Petrides and Pandya, 1988). In addition, it is through the superior occipitofrontal fascicle that the prefrontal cortex maintains strong connections with limbic and paralimbic association areas, particularly the cingulate gyrus, retrosplenial cortex, parahippocampal area, and the presubiculum (Nieuwenhuys et al., 1988) (but see Ture et al., 1997).

The finding of a highly heritable medial frontal cortex bilaterally, Heschl's gyrus bilaterally, and left postcentral gyrus is in agreement with previous reports (Thompson et al., 2001; Wright et al., 2002), underscoring the relevance of these brain areas when searching for genes influencing brain structure and function. Additionally, we found the superior frontal cortex bilaterally, the occipital (striate and extrastriate) cortex in the left hemisphere, the left posterior cingulate, the amygdala bilaterally, and the right parahippocampal gyrus to be highly heritable. Thus, it seems that the individual variation in morphology of areas involved in attention, language, visual, and emotional processing, as well as in sensorimotor processing are strongly genetically influenced. Unique environmental factors influenced vast GM and WM areas surrounding the lateral ventricles (up to 0.50). This finding coincides with the significant environmental influences on lateral ventricle volume [common (0.58) and unique (0.42) with no significant contributions of genes] that we reported previously in this sample (Baaré et al., 2001).

However, we also found differences in our study compared with previous findings in healthy adult twin samples. We did not find significant heritability for Broca's language area (Thompson et al., 2001), ventrolateral prefrontal cortex, and anterior cingulate gyrus (Wright et al., 2002). There may be several reasons for these differences. First, the transformation procedures used in our study (linear followed by nonlinear warping) differed from the transformation procedures used in the other two studies, which applied cortical brain mapping (Thompson et al., 2001) and linear warping (Wright et al., 2002) procedures. Second, our structural equation modeling procedure with correction for multiple comparisons based on random field theory differed from the approach used in the other two studies. Thompson et al. (2001) applied twice the difference between MZ and DZ intraclass correlation coefficients and a permutation analysis to correct for multiple comparisons (Thompson et al., 2001). Wright et al. (2002) adopted a modeling strategy similar to the one used in this study but did not correct for multiple comparisons. Thus, although our method of structural equation modeling, allowing for separate estimates of additive genetic, common environmental, and unique environmental influences, can be considered an optimal model for genetic studies in twin pairs, the conservative approach of using the random field theory for correction for multiple comparisons in these dependent samples may have prevented some areas from reaching significant heritability but allowed the areas with significant heritability to be considered reliable.

A common genetic origin of brain structure shared with PIQ was found to be maximal in the superior occipitofrontal fascicle bilaterally, left optic radiation, right medial frontal cortex, and right parahippocampal gyrus and was still considerable in the left occipital cortex and left corpus callosum. The observed (phenotypic) correlations of up to 0.35 and the genetic correlations of up to 0.39 were higher than those found for the observed and genet-
Gray matter influences focal GM and WM densities and the set of genes that influence VIQ/PIQ. The extent of the overlap is reflected by the magnitude of the genetic correlation ($r_g$), which reflects the correlation between the set of genes that influences focal GM and WM densities and the set of genes that influences VIQ/PIQ. Note that, for illustration purposes, positive cross-correlations as shown here were not thresholded for significance. By definition, the cross-correlations in voxels that were not significant were set to zero. The observed correlation ($r_p$) reflects the correlation that is found between GM and WM density and VIQ/PIQ in individuals. The cross-trait/cross-twin correlations show the correlations of focal GM and WM densities of a twin with the intelligence quotient of his/her cotwin and vice versa for MZ and DZ twin pairs (or sibling pairs) separately. If the correlation between GM and WM density of a twin and VIQ or PIQ in the cotwin is larger in MZ twins than in DZ twins, this indicates that the genes influencing GM and WM density partly overlap with the genes that influence VIQ/PIQ. The extent of the overlap is reflected by the magnitude of the genetic correlation ($r_g$). $r_g$ reflects the correlation between the set of genes that influences focal GM and WM density and the set of genes that influences VIQ/PIQ.

Table 3. Observed (phenotypic) correlations, cross-trait/cross-twin correlations, and genetic correlations.

<table>
<thead>
<tr>
<th>Left hemisphere, IQ ($x, y, z$)</th>
<th>Observed ($r_p$)</th>
<th>MZ DZ Genetic ($r_g$)</th>
<th>Cross-trait/cross-twin</th>
<th>Correlation</th>
<th>Cross-trait/cross-twin</th>
<th>Correlation</th>
<th>Right hemisphere, IQ ($x, y, z$)</th>
<th>Observed ($r_p$)</th>
<th>MZ DZ Genetic ($r_g$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gray matter</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Gray matter</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Occipital cortex, PIQ ($-21, -62, 13$)</td>
<td>0.17</td>
<td>0.08</td>
<td>-0.06</td>
<td>0.23</td>
<td></td>
<td></td>
<td>Medial frontal, PIQ ($41, 22, 43$)</td>
<td>0.25</td>
<td>0.32</td>
</tr>
<tr>
<td>Superior occipitofrontal fascicle, PIQ ($-21, -24, 35$)</td>
<td>0.17</td>
<td>0.18</td>
<td>0.04</td>
<td>0.24</td>
<td></td>
<td></td>
<td>Parahippocampal, V1Q ($23, -38, -7$)</td>
<td>0.11</td>
<td>0.15</td>
</tr>
<tr>
<td>Corpus callosum, PIQ ($-15, 3, 33$)</td>
<td>0.18</td>
<td>0.17</td>
<td>0.17</td>
<td>0.22</td>
<td></td>
<td></td>
<td>Superior occipitofrontal fascicle, PIQ ($21, -29, 31$)</td>
<td>0.35</td>
<td>0.25</td>
</tr>
<tr>
<td>Optic radiation, PIQ ($-35, -48, 17$)</td>
<td>0.26</td>
<td>0.28</td>
<td>0.01</td>
<td>0.39</td>
<td></td>
<td></td>
<td>Corpus callosum, V1Q ($15, 3, 37$)</td>
<td>0.14</td>
<td>0.13</td>
</tr>
</tbody>
</table>

The observed correlation ($r_p$) reflects the correlation that is found between GM and WM density and VIQ/PIQ in individuals. The cross-trait/cross-twin correlations show the correlations of focal GM and WM densities of a twin with the intelligence quotient of his/her cotwin and vice versa for MZ and DZ twin pairs (or sibling pairs) separately. If the correlation between GM and WM density of a twin and VIQ or PIQ in the cotwin is larger in MZ twins than in DZ twins, this indicates that the genes influencing GM and WM density partly overlap with the genes that influence VIQ/PIQ. The extent of the overlap is reflected by the magnitude of the genetic correlation ($r_g$). $r_g$ reflects the correlation between the set of genes that influences focal GM and WM density and the set of genes that influences VIQ/PIQ.

The finding of a common genetic origin of frontal and occipital GM with intelligence is corroborated by findings from two magnetic resonance imaging studies in singletons, in which several brain areas were found to be correlated with intelligence (Frangou et al., 2004; Haier et al., 2004). Interestingly, frontal and occipital GM has been associated with general fluid intelligence (Spearman’s $g$) in a functional positron emission tomography study in singletons (Duncan et al., 2000). Moreover, in a functional magnetic resonance imaging study, greater fluid intelligence was associated with larger event-related neural activity in the prefrontal cortex, quite comparable with that found to be genetically determined in our study (Gray et al., 2003). A role for the superior occipitofrontal fascicle in (nonverbal) intelligence is corroborated by a study suggesting this pathway to be implicated in spatial attention and visuospatial functioning (Filley, 2001). We found evidence that (nonverbal) intelligence shares common genes with frontal and occipital GM areas, in concert with the connecting superior occipitofrontal fascicle and interhemispheric corpus callosum.

Most focal brain areas that were primarily influenced by genes were represented on both sides of the brain. Thus, both the left- and right-hemisphere representations of anatomical brain regions shared the extent to which their individual differences in GM and WM density were genetically determined. Indeed, GM was found to be a good predictor of the density of the homotopic region in the contralateral hemisphere, with the striking exception of the primary visual cortex (Mechelli et al., 2005). Our findings suggest that the compatibility of GM density of several homotopic areas in the contralateral hemisphere is genetically determined. Moreover, our findings suggest that individual variation in GM density of the primary visual cortex is genetically mediated.

In this study, we measured the extent of genetic contributions and not the influence of individual genes on brain morphology. However, our findings are supported by a study in which a functional variation in the val66met locus in the 5’ prodomain of brain-derived neurotrophic factor (BDNF) was found to influence brain morphology in normal humans. Individuals who were met-BDNF carriers compared with individuals who were val/val-BDNF carriers had reduced volumes of the medial frontal GM bilaterally and the right hippocampus, which has close anatomical connections with the parahippocampal gyrus (Pezawas et al., 2004). Considering that the BDNF gene has a role in human learning and memory, this gene may be one of the candidates.
implicated in a positive association between the medial frontal and medial temporal cortices and intelligence that is attributable to common genes.

In conclusion, we found that specific focal GM and WM areas in the human brain are highly heritable. Moreover, some of these GM and WM areas, including the superior occipitofrontal fascicule, corpus callosum, and medial frontal and occipital cortices, share common genes with intelligence, forming a heritable neural network in the human brain that is involved in intelligence.

References


