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ORIGINAL ARTICLE

# Human Cortical Thickness Organized into Geneticallydetermined Communities across Spatial Resolutions

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## Abstract

The cerebral cortex may be organized into anatomical genetic modules, communities of brain regions with shared genetic influences via pleiotropy. Such modules could represent novel phenotypes amenable to large-scale gene discovery. This modular structure was investigated with network analysis of in vivo MRI of extended pedigrees, revealing a "multiscale" structure where smaller and larger modules exist simultaneously and in partially overlapping fashion across spatial scales, in contrast to prior work suggesting a specific number of cortical thickness modules. Inter-regional genetic correlations, gene co-expression patterns and computational models indicate that two simple organizational principles account for a large proportion of the apparent complexity in the network of genetic correlations. First, regions are strongly genetically correlated with their homologs in the opposite cerebral hemisphere. Second, regions are strongly genetically correlated with nearby regions in the same hemisphere, with an initial steep decrease in genetic correlation with anatomical distance, followed by a more gradual decline. Understanding underlying organizational principles of genetic influence is a critical step towards a mechanistic model of how specific genes influence brain anatomy and mediate neuropsychiatric risk.

Key words: gene expression, genetic correlations, modularity, network analysis, structural MRI

# Introduction

A primary aim of imaging genomics is to identify specific genes that influence neuroanatomical variation (Glahn et al. 2007). As genetic influences on human brain anatomy are profound (Pennington et al. 2000; Posthuma et al. 2000; Baaré et al. 2001; Schmitt et al. 2007a), it is likely that specific genes influence individual differences observed in morphometric analyses. Furthermore, these genes are strong candidate genes for illnesses associated with neuropathology (Glahn et al. 2014). Heritability, the proportion of population variability due to genetic factors, has been quantified for univariate brain traits such as volume,

thickness or surface area at whole-brain (Bartley et al. 1997), lobar (Geschwind et al. 2002), gyral (Peper et al. 2007) and subgyral resolutions (Stein et al. 2010). Typically, these analyses involve parcellating an MRI image into anatomically distinct regions (e.g., Fischl et al. 2004b). Large-scale genome wide association studies (GWAS) have been conducted on these parcellated traits, providing a number of replicable loci (Bis et al. 2012; Stein et al. 2012; Hibar et al. 2015; 2017). Additionally, single-gene mutations cause specific cortical dysmorphologies (Gaitanis and Walsh 2004; Guerrini and Marini 2006). However, specific genes that influence normal anatomical variation have yet to be identified.

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This limited progress could reflect the complexity of genetic influences. Individual brain traits are thought to be influenced by a complex relationship of environmental factors, common variants of small effects and rare variants of large effects (Thompson et al. 2013). While it is theoretically possible that the genetic influences on different parcellated brain regions are independent of one another, the evidence for such genetic autonomy is limited. Rather, multiple morphometrically distinct brain regions have common genetic roots (Baaré et al. 2001; Schmitt et al. 2007b, 2010; Glahn et al. 2015; McKay et al. 2015), suggesting pleiotropic effects whereby the same genes influence distributed neuroanatomical networks. Ontogenetically, it is possible that shared genetic variance results when neurons that comprise 2 regions derive from similar germinal zones at similar developmental periods. Genetically unique brain systems could provide novel phenotypes amenable to large-scale genome-wide search, which would constitute a fundamental advance in the genetics of human neuroanatomy. The application of formal network analysis (Bullmore and Sporns 2009; Sporns 2010) based upon genetic correlation provides a method for delineating neuroanatomic networks defined by common genetic factors. Therefore, we used graph theoretical techniques to investigate any potential genetic network architecture of the adult human brain.

In a series of recent studies, Chen and colleagues (Chen et al. 2011a, 2012, 2013), applied various network methods in ~200 twin pairs to investigate patterns of inter-regional genetic correlations, reporting the genetic modularity of neuroanatomic networks. More specifically, Chen and colleagues identified 12 specific modules (or clusters). Although cortical surface area and cortical thickness were not genetically correlated (Panizzon et al. 2009; Winkler et al. 2010), these 13 fundamental modules were largely overlapping between these morphometric indices. Identified modules were largely bilaterally symmetric (left and right hemispheres were mirror images) and spatially contiguous (within hemispheres modules were composed of anatomically adjacent regions). Given that brain networks across multiple modalities tend to have a scale-free organizational structure (Meunier et al. 2009; Bassett et al. 2010), evidence for a specific modular resolution (a single "natural" number of modules) is particularly striking and has profound implications for studies of the regional specificity of brain function, pathophysiology and gene discovery. Unfortunately, the modular networks identified by Chen and colleagues have yet to be replicated in an independent sample.

The present study extends this prior work on the genetics of brain cortical architecture in several ways. First, this study is one of the largest studies to date of genetic correlations in cortical thickness between brain regions, using a sample of ~1400 individuals in large, extended pedigrees. Second, we use network theory and machine learning techniques to explore whether the brain is organized into a "natural" number of genetic correlation modules. Third, we investigate the relationship between genetic correlations and the anatomical distance between brain regions, as spatial constraints are known to influence network properties (Vértes et al. 2012). Finally, we compare the imaging-genetic findings with correlations in gene expression from postmortem microarray studies (Hawrylycz et al. 2012). Our findings suggest a multiscale, multiresolution modular structure consistent with a hierarchical organization, where smaller components are iteratively connected into larger components and systems. Relatively simple organizational principles, including contralateral homology and greater genetic overlap with greater anatomical proximity, underlie much of the apparent complexity observed in the network of genetic correlations.

## **Methods**

# **Imaging Acquisition and Processing**

Since 2006, the Genetics of Brain Structure and Function study (GOBS), has recruited randomly ascertained extended pedigrees of Mexican American descent living in San Antonio (McKay et al. 2014). High-resolution structural MRI scans were acquired on a Siemens 3T TIM Trio at UTHSCSA. The sequence was composed of multiple T1-weighted 3D turbo-flash sequences with an adiabatic inversion contrast pulse (0.8 mm isotropic voxels, TE 3 ms, TR 2100 ms, TI 785 ms, flip angle 13°, FOV 200 mm). The sample included 1443 individuals (836 female) with mean age of 40.7 years (SD = 15.5, range = 18-85).

Image processing used FreeSurfer version 5.3 (Dale et al. 1999; Fischl et al. 1999; 2001; Fischl et al. 2004a, 2004b; Desikan et al. 2006). Briefly, after normalization and skull stripping, whitematter voxels were identified based on intensity and covered in a topologically corrected, tessellated mesh. This gray-white surface was then expanded to fit the pial surface. Surfaces were examined manually and edited if necessary. Cortical curvature was used to register surfaces to a common atlas (Desikan atlas) and identify 68 regions interest (Desikan et al. 2006). Cortical thickness was calculated as the distance between the gray-white surface and the pial surface at each vertex.

A replication dataset was also analyzed, using publically available data from the human connectome project (HCP; http://www.humanconnectome.org/), which comprised MRI data from 1113 individuals (606 female) from 457 unique families (including 170 dizygotic twins, 286 monozygotic twins, 576 non-twin siblings, and 25 non-sibling familial relations) with mean age 28.8 years (SD = 3.7, range = 22-37). As previously described (Glasser et al. 2013; Van Essen et al. 2013), T1weighted and T2-weighted structural images were acquired on a 3T Siemens Skyra employing a 32-channel head coil. T1 scans used a magnetization prepared rapid gradient echo sequence (0.7 mm isotropic voxels, TE 2.14 ms, TR 2240 ms, TI 1000 ms, flip angle  $8^{\circ}$ , ES 7.6 ms, GRAPPA = 2.). T2 scans used a variable flip angle turbo spin-echo sequence (0.7 mm isotropic voxels, TE 565 ms, TR 3200 ms,  $BW = 744 \, Hz$  per pixel, no fat suppression pulse, GRAPPA = 2). Preprocessing was performed using version 3 of the HCP pipeline (Glasser et al. 2013), including coregistration of T1 and T2 scans, B1 (bias field) correction, and segmentation and surface reconstruction using FreeSurfer version 5.3-HCP to estimate cortical thickness in 68 regions interest (Desikan et al. 2006).

## **Genetic Correlation Network**

Variance-components genetic analysis was performed to differentiate genetic and environmental contributions to the phenotypic covariance between cortical thicknesses of different brain regions (Williams et al. 1999). For an individual trait such as the cortical thickness of one region, heritability is estimated using the polygenic model (Almasy and Blangero 1998; Lynch and Walsh 1998)

$$y = X\beta + g + e \tag{1}$$

where  $\mathbf{y}$  is the trait vector,  $\mathbf{X}$  is a matrix of covariates,  $\mathbf{\beta}$  is a vector of coefficients, g is a vector reflecting the random additive effect of genetics, and e is a vector of residuals (which includes environmental effects not modeled). The vector e follows a multivariate normal distribution with zero mean and variance-covariance matrix  $\sigma_q^2 \mathbf{M}$ , where  $\sigma_q^2$  is a scalar reflecting the additive genetic variance, and M is twice the kinship matrix (quantifying the genetic relationships between subjects). The vector e follows a multivariate normal distribution with zero mean and variancecovariance matrix  $\sigma_e^2 \mathbf{I}$ , where  $\sigma_e^2$  is the residual variance, and  $\mathbf{I}$  is the identity matrix. The vector **y** is distributed as  $y \sim N(\mathbf{X}\boldsymbol{\beta}, \sigma_a^2 \mathbf{M} +$  $\sigma_e^2 \mathbf{I}$ ), where  $\beta$ ,  $\sigma_g^2$ , and  $\sigma_e^2$  are free parameters. The heritability  $(h^2)$  of the trait is given by  $h^2 = \sigma_q^2/(\sigma_q^2 + \sigma_e^2)$ . Maximum likelihood estimation was performed with SOLAR software (www. solar-eclipse-genetics.org) (Almasy and Blangero 1998).

The univariate polygenic model described above can be extended to estimate the genetic and environmental variances of 2 traits, A and B, as well as the genetic and environmental correlations between them, using the bivariate polygenic model

$$\mathbf{y}_{\mathbf{A}\mathbf{B}} = \mathbf{X}_{\mathbf{A}\mathbf{B}}\mathbf{\beta}_{\mathbf{A}\mathbf{B}} + \mathbf{g}_{\mathbf{A}\mathbf{B}} + \mathbf{e}_{\mathbf{A}\mathbf{B}} \tag{2}$$

where  $\mathbf{y}_{\!_{\mathbf{AB}}}$  is a concatenated vector containing both traits. The variance–covariance matrix of  $\boldsymbol{g}_{AB}$  is  $\boldsymbol{G}\otimes\boldsymbol{M},$  where  $\otimes$  is the Kronecker product, and  ${\bf G}$  is a 2-by-2 matrix whose diagonal elements are the genetic variances of A and B ( $\sigma_{qA}^2$  and  $\sigma_{eB}^2$ , respectively) and whose off-diagonal elements are the covariance between the genetic effects  $(\sigma_{qAB}^2)$ . The variance–covariance matrix of  $e_{AB}$  is  $\textbf{E} \otimes \textbf{I}$ , where E contains the residual variances and covariance ( $\sigma_{eA}^2$ ,  $\sigma_{eB}^2$ , and  $\sigma_{eAB}^2$ ). After maximum likelihood estimation with SOLAR, the genetic correlation is calculated as  $\rho_{g} = \sigma_{gAB}^{2} / \sqrt{\sigma_{gA}^{2} \cdot \sigma_{gB}^{2}}$ . Age, age<sup>2</sup>, sex, and their interactions, as well as intracranial volume, were included as covariates in all analyses.

Genetic correlations were calculated between all 2278 pairs of macro-anatomic brain regions. The absolute value of these  $\rho_q$  formed the elements of a 68  $\times$  68 association matrix, and weighted network models were constructed by thresholding these matrices at different edge densities (the proportion of elements of the association matrix included in the network as edges) including minimum spanning trees to ensure all networks were connected (Alexander-Bloch et al. 2010). Networks were constructed at each density between 0 and 0.2 at 0.001 intervals. Note that at this range of connection densities, the magnitude of positive correlations always exceeded that of negative correlations, so the results were robust to taking or not taking the absolute value of  $\rho_g$ .

The modularity, Q, of these weighted networks was estimated as  $Q = \frac{1}{2m} \sum_{i \neq j} (A_{ij} - P_{ij}) \delta(M_i, M_j)$ , where m is the total weight of the edges in the network;  $A_{ij} = \rho_G$  if there is an edge between i and j (otherwise  $A_{ij} = 0$ ); the delta function  $\delta = 1$  if i and j are in the same module M (otherwise  $\delta = 0$ ); and  $P_{ij} = \frac{k_i k_j}{2m} \text{ is the expected weight of an edge between } i \text{ and } j,$  where  $k_i$  is the total weight of i's edges. At each density between 0 and 0.2, the optimal modular partition was estimated using Newman's algorithm (Clauset et al. 2004).

Representative consensus partitions across densities were then calculated by successively matching partitions by solving the linear assignment problem (Dimitriadou et al. 2002) in order of their average normalized mutual information with other partitions (Strehl and Ghosh 2002). Network analysis was performed in R using iGraph (Csardi and Nepusz 2006), Clue (Hornik 2005) and Cluster (Maechler et al. 2012) software packages. All networks were compared with randomly generated networks with same number of nodes, connection densities and degree distributions (Viger and Latapy 2005).

Two additional analyses were performed to specifically query the modular structure across scales or resolutions.

Newman's method may be biased against smaller modules with fewer nodes (Fortunato and Barthélemy 2007), which was addressed by the addition of a resolution or scale parameter,  $\gamma$ , to the modularity quality function which was optimized alternatively with simulated annealing (Reichardt and Bornholdt 2004) and with the generalized Louvain method (Blondel et al. 2008; De Meo et al. 2011) (see Supplementary Materials). A plateau in the optimal number of modules across a range of values for this parameter, or a peak in the similarity of the modular structure across multiple instantiations of the algorithm, may indicate a more natural number of modules or specific spatial scale (Arenas et al. 2008; Betzel et al. 2017) as opposed to there being no preference for a specific resolution or scale. In addition, partitioning around medoids (PAM) clustering was performed on the genetic correlation matrix. When queried across a range of possible clusters (the number of clusters, k, is an input parameter for this clustering method), a sharp peak in a distinct quality function such as the silhouette width can indicate a more or less "natural" split into a number of homogenous clusters (Kaufman and Rousseeuw 1987). Alternatively, a lack of such a peak can be interpreted as evidence of a lack of a specific and distinct number of clusters.

The relationship between anatomical distance and  $ho_g$  was estimated using general linear models in R statistics (R Core Team 2016). For polynomial models

$$\boldsymbol{\rho_{\mathrm{g}}} = \beta_0 + \beta_1 \mathbf{d} + \beta_2 \mathbf{d}^2 \cdots + \beta_n \mathbf{d}^{\mathbf{n}} + \mathbf{e} \tag{3}$$

where d is a vector of the Euclidean between all pairs of regions (distance between the centroids of the regions on the pial surface) and e is a vector of residuals. This model was fit separately for intra- and inter-hemispheric pairs of regions, where for interhemispheric regions, distance was estimated as the distance from a region's contralateral homolog. Linear, quadratic, cubic and higher order polynomial models were compared using likelihood ratio tests and Akaike Information Criterion (Venables and Ripley 2002). Exponential decay models

$$\rho_{\mathbf{q}} = \beta_0 \exp(-\beta_1 \mathbf{d}) + \mathbf{e} \tag{4}$$

were compared with polynomial models using the Akaike Information Criterion. The significance of the relationship with distance was also tested with Mantel permutation tests.

# **Gene-Expression Networks**

Gene-expression networks were derived from publicly available microarray data from the Allen Human Brain Atlas (http:// human.brain-map.org/) (Hawrylycz et al. 2012). Postmortem brain tissue was collected from 6 individuals, with ~500 samples collected from each hemisphere; left hemisphere data was collected in all 6 subjects, while right hemisphere data was collected in 2 subjects. Expression levels for ~60 000 RNA probes were collected for each sample. The data were averaged across probes corresponding to the same gene, resulting in 20739 genes from 3702 samples. MNI coordinates corresponding to each sample were matched to the FreeSurfer average cortical surface, allowing for comparisons with the imaging-genetics data. Only the 1745 cortical samples were included in subsequent analysis, with the subcortical samples excluded for consistency with the analysis of cortical thickness in the brain imaging data. Data within each gene were standardized, and gene expression correlations (r<sub>GE</sub>) were calculated using Pearson's r between every pair of cortical samples, providing a measurement of the similarity in the samples' gene-expression profiles.

The within-module and between-module  $r_{GE}$  was calculated by assigning each cortical sample to its nearest  $ho_{
m g}$  module. The within-module and between-module  $r_{GE}$  was calculated for each subject individually and then averaged across subjects. The significance of within module  $r_{GE}$  was assessed by comparing these data to permutations where the module membership of each region was randomized. The anatomical distance between brain samples was estimated using the Euclidean distance between the MNI coordinates of each sample. Contralateral homology was inferred from the anatomical labels for the two subjects with bilateral data. Differential expression between cortical modules for each gene was calculated using lme4 in R (Bates et al. 2014), using subject and region as random effects and module as a fixed effect. P-values were obtained by likelihood ratio tests of the full model including module membership compared with the reduced model without module membership. The results of this test were entered into a gene ontology (GO) analysis using TopGO in R (Alexa et al. 2006) with Entrez gene annotations and Kolmogorov-Smirnov test of enrichment. FDR correction for multiple comparisons was performed on the results of the enrichment test (Benjamini et al. 2006).

## Computational Model

We used a simple computational model to investigate whether the modularity of the  $\rho_a$  network was related to or driven by the constraints of anatomical distance and contralateral homology, by generating null networks where these properties were preserved but wiring was otherwise generated at random. For the genetic correlation network, the relationship between anatomical distance and  $ho_g$  was estimated with polynomial models as described above. As described in the Results, the best-fit model was a quadratic polynomial model for intrahemispheric pairs of regions, and a cubic polynomial model for interhemispheric pairs of regions. For each pair of regions, the predicted  $\rho_{q}$  from these models,  $\rho_{qP}$ , was an estimate of the genetic correlation expected based on anatomical distance constraints and contralateral homology alone. The final model of genetic correlation,  $\rho_{\rm gM}$  , also included a stochastic element  $\theta$ 

$$\rho_{g\mathbf{M}} = \rho_{g\mathbf{P}} + \theta \tag{5}$$

where  $\theta$  was a vector of noise, derived by adding or subtracting 0.05 (equivalent to 5% of the highest magnitude genetic correlation in the real data) from pairs of regions randomly chosen with replacement. The number of pairs of regions chosen was varied such that the stochastic element accounted for 0-50% of the weight of the total genetic correlation matrix (in 1% intervals) in the modeled networks. As this number increases, modeled networks are predicted to increasingly reflect topological characteristics of random networks.

The modular characteristics of these modeled networks were compared with those of the actual  $ho_q$  network to determine whether a given "value" of  $\theta$  (the proportion of the modeled network that was based on the stochastic process) generated model networks with similar modularity as the actual network. For this purpose, the mean modularity was calculated across the range of density thresholds (from 0.0 to 0.2 in 0.001 intervals). Finally, for model networks with the same modularity as the actual network, the within module gene expression correlations,  $r_{GE}$ , were calculated to determine whether the network model could account for the relationship between genetic correlation modules and inter-regional correlations in gene expression profiles.

## Results

#### Genetic Correlations Reveal Modular Structure

The pattern of genetic correlations  $(\rho_a)$  formed a complex, heterogeneous network (Fig. 1A). A greater proportion of the shared variance in cortical thickness between regions was due to shared genetic factors (mean  $\rho_q = 0.48$ , SD = 0.19, range = –0.17–1.0) than to environmental factors (mean  $\rho_e$  = 0.22, SD = 0.13, range = -0.18-0.74, see Supplementary Fig. 1). Although there were some negative genetic correlations, the magnitude of the high positive correlations exceeded those of the negative correlations, so the negative correlations were not included in the networks derived from the inter-regional genetic correlations. This network had significant modular structure (Fig. 1B) compared with randomized, connected networks with the same degree distribution, at every connection density tested (P < 0.01, 1000 random networks per connection density).

In contrast, there was not a significant difference in the optimum number of modules for the  $\rho_q$  network compared with random networks (Fig. 1B). Across graph densities, optimal partitions are found across a wide range in terms of the number of modules, suggesting that there is no single number of modules that satisfactorily represents the structure of the network (Fig. 1B). Supplementary analysis, including silhouette plots of k-medoid (PAM) clusters and the addition of a resolution parameter,  $\gamma$ , to identify modules across spatial scales, further confirmed this multiscale modular structure (see Supplementary Fig. 2).

The modular partitions appeared to largely respect a priori functional boundaries. In particular, medial occipital, dorsallateral prefrontal, ventral-medial prefrontal and the limbic system were relatively well delineated (Fig. 1C). At a coarser resolution (2 and 5 module solutions in Supplementary Fig. 3) the brain was subdivided principally along a dorsal-ventral axis. It is notable that, although the algorithms were in no way constrained to cluster spatially adjacent regions, nor to cluster contralateral homologous regions, modules were largely composed of laterally symmetric, spatially contiguous regions of the cortical surface. Contralateral symmetry has previously been observed in modules derived from functional connectivity (Meunier et al. 2009) and structural covariance (Chen et al. 2011b) (which combines genetic and environmental correlation). Notable exceptions to contralateral symmetry were regions within the inferior frontal gyrus, which in the right hemisphere were located in the dorsallateral (red) and ventral medial (light blue) module; while in the left hemisphere these regions were all located within the dorsallateral module. There was only limited overlap between pairs of regions with high genetic correlations and those with relatively high environmental correlations (see Supplementary Fig. 1).

Quantitatively, genetic modules were compared with the recent multimodal parcellation generated from HCP data (Glasser et al. 2016). Although the limitations of this kind of comparison should be noted (Alexander-Bloch et al. 2017; see Supplementary Materials), there appeared to be a high correspondence, such that in each hemisphere the 180 HCP parcels tended to lie within genetic modules. Using the 8-module partition for example, nearly half of the HCP-parcels were >90% within a single genetic module, respectively (mean = 84% within a single genetic module, median 89%, range = 36-100%).

# The Influence Anatomical Distance and Contralateral Homology

As suggested by the modular structure, the genetic correlations between contralateral homologs were greater on average than

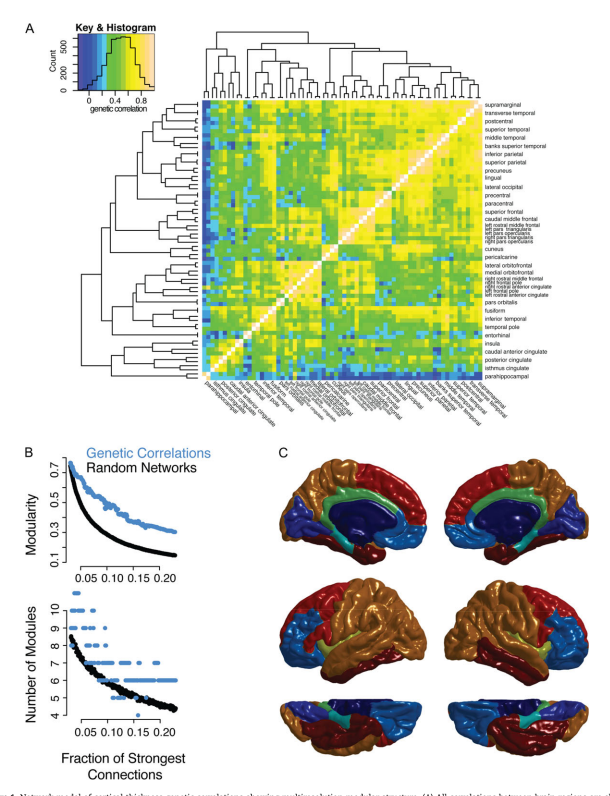


Figure 1. Network model of cortical thickness genetic correlations showing multiresolution modular structure. (A) All correlations between brain regions are shown with color scale indicating strength of genetic correlation. The regions are organized by complete linkage hierarchical clustering, with the dendrogram shown to the left and to the top of the heat map. Names of regions are shown to the right and to the bottom of the heat map; hemisphere (left or right) is specified only for regions that were not clustered such that they are adjacent to their contralateral homolog. Color key is shown to the top left, as well as a histogram showing the number of pairs of regions (count) with genetic correlations at each strength. (B) Network models (graphs) were constructed at each connection density from 0.0 to 0.2 in 0.001 intervals, where the connections density refers to the fraction of the strongest connections that are included in the graph. The genetic correlation networks are shown in addition to random control networks (see Methods). The top plot shows modularity, a measure of the strength of the community structure of the network, which is significantly higher in the genetic correlation network compared with the random network. The bottom plot shows the optimal number of modules at each density. (C) An illustration of the consensus partition (modular community structure) with 8 cortical modules (see Supplementary Fig. 3 for illustrations of the partition at different resolutions).

the genetic correlations between other regions (mean genetic correlation with contralateral homolog,  $\rho_q = 0.94$ , SD = 0.06, range = 0.74-1.0). Of all 68 regions, only 5 regions had higher genetic correlations with any other region than with their homolog in the opposite hemisphere; exceptions included canonical language areas with known functional asymmetry (left bank of the superior temporal sulcus, left pars opercularis, left pars triangularis) as well as left rostral middle frontal gyrus and right caudal anterior cingulate. This reflects a general pattern where the genetic correlations with contralateral homologs were relatively lower in the cingulate and in inferior prefrontal regions compared with the rest of the cortex (Fig. 2C). The magnitude of environmental correlations between contralateral homologs was lower than that of genetic correlations, both in absolute terms and relative to environmental correlations with non-homologous regions (see Supplementary Fig. 1).

Genetic correlations were strongly influenced by the anatomical distance between brain regions (Fig. 2A, B). For intrahemispheric genetic correlations, this relationship was approximated by a quadratic polynomial model (residual difference in sum of squares compared with a linear model = 0.67, Chi square test (1 df) P < 1e-5). There was no evidence for a significant cubic relationship with intrahemispheric anatomical distance (residual difference in sum of squares compared with quadratic model = 0.07, Chi square test (1 df) P = 0.12). The quadratic model also outperformed an exponential decay model (adjusted  $r^2$ -quadratic = 0.15; adjusted  $r^2$ -quadratic exponential = 0.04; Akaike information criterion quadratic = -737; Akaike information criterion exponential = 2097).

The relationship between genetic correlations and anatomical distance was regionally heterogeneous. This relationship also diverged from that between anatomical distance and the environmental correlations between regions. Although there was a negative relationship between anatomical distance and environmental correlations, this relationship largely plateaued around 4cm, while the relationship with genetic correlations extended to

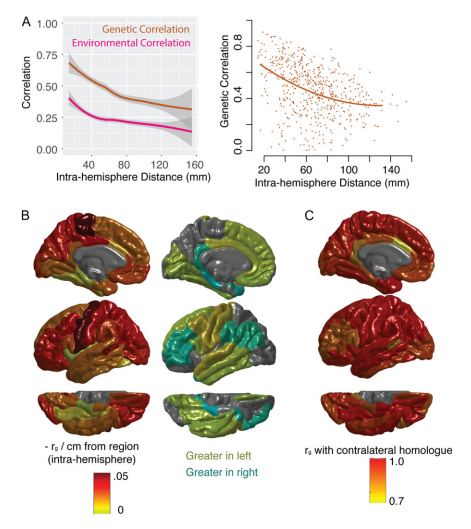


Figure 2. The relationship between genetic correlation  $(\rho_g)$ , anatomical distance and contralateral homology. (A) The left-hand plot shows the relationship between anatomical distance and phenotypic (total) correlation, as well as genetic correlation ( $\rho_g$ , the genetic component of the phenotypic correlation), and environmental correlation (the non-genetic component of the phenotypic correlation). Lines represent smoothed local averages, with standard error silhouettes (ggplot2 package in R). The right-hand plot focuses on the genetic correlation  $(g_g)$ , showing the fit of a quadratic model of the relationship with distance, which is a better fit than either a linear model or an exponential decay model (see Results). These plots show correlations within the left hemisphere only. (B) For each region, the linear relationship between  $\rho_q$  between that region and every other region was estimated. At the regional level, the model fits were not improved by adding quadratic components. Color scheme represents the estimated decrease in  $\rho_g$  per cm from the region of origin. The hemispheres were averaged for the left-hand plot. The right-hand plot whether the slope of the linear relationship with distance was greater in the left hemisphere (yellows) or right hemisphere (aqua), for each region. (C) The  $\rho_g$  between each region and its contralateral homolog in the opposite hemisphere.

longer distances (Fig. 2A). When the relationship of  $\rho_q$  with distance was estimated for each region separately, lateral and dorsal aspects of the cortex appeared to have stronger negative relationships with distance; relatively weaker relationships with distance existed in ventral and temporal regions (Fig. 2B). Although largely convergent between hemispheres, certain regions diverged from their contralateral homologs in terms of the strength of there with distance, including canonical language areas (Fig. 2B).

# Replication Data

A similar network structure was observed in the  $ho_q$  network derived from the HCP data. The network demonstrated significant modularity at every connection density, but did not demonstrate a single optimal number of modules (Fig. 5A). The exact modular partitions derived for the GOBS and HCP data were similar but not identical. For example, prefrontal cortical regions that belonged to different modules in the GOBS data (Fig. 1C) belong to the same module in the HCP data (Fig. 5D). Similar relationships with anatomical distance and contralateral homology were also observed in the HCP data. The HCP genetic correlations were strongly influence by the anatomical distance between brain regions (Fig. 5B). In addition, the HCP genetic correlations between contralateral homologs were greater on average than the genetic correlations between other regions (Fig. 5C), recapitulating the results from the GOBS data.

# Gene Expression, Genetic Correlation

Correlations in gene expression,  $r_{GE}$ , were higher within  $\rho_a$ modules as opposed to between  $\rho_a$  modules, despite the fact that these modules were derived independently from the gene expression data (Fig. 3A, B). Compared with a null model where samples were randomly assigned to modules, the withinmodule genetic correlation was significantly higher when regions were assigned to their actual module (P < 0.001, 1000 permutations). Particularly strong correlations in gene expression profiles occurred within the occipital module (dark blue) and within the dorsal prefrontal module (red), although the prefrontal module had higher between-module r<sub>GE</sub> with other modules than did the occipital module (Fig. 3B).

Similar to the  $\rho_q$  networks,  $r_{\rm GE}$  networks also revealed the influence of anatomical distance and contralateral homology. For the 2/6 subjects with gene expression data from both cortical hemispheres, the  $r_{GE}$  between a brain region and its contralateral homolog was greater than the average  $r_{\rm GE}$  between that

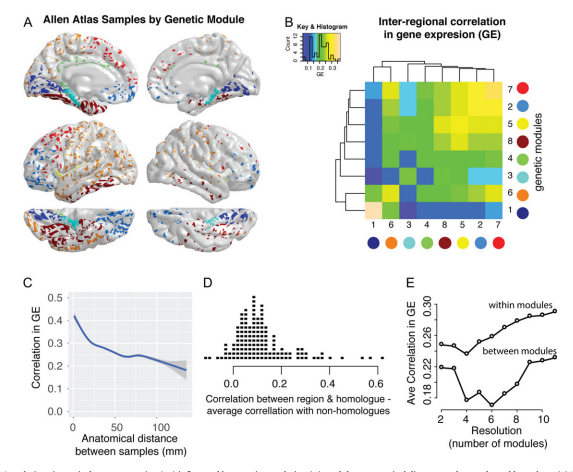


Figure 3. Correlations in cortical gene expression ( $r_{GE}$ ) influenced by genetic correlation ( $\rho_g$ ) modules, anatomical distance and contralateral homology. (A) Location of cortical samples available for 6 subjects in Allen Atlas, color-coded based on  $\rho_g$  module (as per Fig. 1C). (B) The intersample correlations in expression pattern across genes were calculated for each pair of samples (within subjects only). The heat map illustrates the average within-module rGE (diagonal elements) and the average between-module  $r_{GE}$  (off-diagonal elements). (C) The relationship between anatomical distance and  $r_{GE}$  (smoothed local average with standard error silhouettes). (D) For each of the FreeSurfer regions (for subjects with samples from both left and right hemispheres), the strip chart shows the difference between r<sub>GE</sub> with the contralateral homolog and the average  $r_{\rm GE}$  with non-homologous regions. (E) The correlation in gene expression patterns across genes across a range of modular resolutions from 2 to 11 genetic correlation modules, showing the average within-module and the average between module rGE across all modules.

region and other brain regions in >90% of cases (Fig. 3D). Comparable to the  $\rho_q$  data, a biphasic relationship also existed between  $r_{GE}$  and anatomical distance, with a steep initial decrease in correlation strength followed by a more gradual decrease at longer anatomical distances (Fig. 3C).

GO analysis showed that the differential expression of genes between cortical genetic modules were linked to plausible biological processes. In total, 66 GO terms survived correction for multiple comparison (FDR-corrected P < 0.05), including nervous system development, brain development, telencephalon development, regulation of stem cell proliferation, chromatin regulation and protein targeting. Supplementary Figure 4 shows the top 5 GO terms and their hierarchical relationships with other biological processes, and Supplementary Table 1 shows all 66 significant GO terms.

The elevated  $r_{GE}$ , within—as opposed to between— $\rho_q$  modules was consistent across a range of modular resolutions (Fig. 3E). For the consensus modular structures from n = 2 (33) regions per module on average) to n = 11 (6 regions per module on average), the average within-module  $r_{GE}$  (mean across resolutions = 0.264, SD = 0.019) always exceeded the average between-module (mean across resolutions = 0.204, SD = 0.023).

## Modeling Genetic-correlation Networks

The constraints of anatomical distance and contralateral homology on genetic correlations appeared to be directly related to the modularity of this network. Models were constructed where the genetic correlation between regions was predicted by these relationships alone, and these models adequately predicted the modularity of the actual  $\rho_a$  network (Fig. 4A,B). As noted above, quadratic models were appropriate for the intrahemispheric relationship with distance, while cubic models were necessary to accurately model the interhemispheric relationship with distance because of the disproportionately high correlations between contralateral homologs.

The modules predicted by the computational model also had significantly elevated within-module  $r_{GE}$ , suggesting that the influence of distance and contralateral homology may drive the observed convergence between  $r_{GE}$  and  $\rho_a$  network organization (Fig. 4C). Importantly, the model's success in predicting within-module  $r_{GE}$  was not due entirely to samples within the same FreeSurfer anatomical region. For networks that respected this cortical parcellation but whose between-region connections were randomized, the within-module  $r_{\text{GE}}$  was

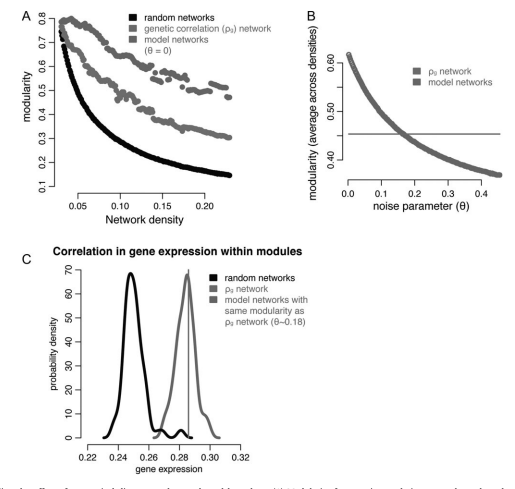


Figure 4. Modeling the effect of anatomical distance and contralateral homology. (A) Modularity for genetic correlation networks and random networks as per Figure 1B. For model networks, inter-regional correlation is based on what would be predicted based on the statistical relationship with anatomical distance within and between hemispheres (Fig. 2) and a noise parameter  $\theta$  ( $\theta$  = 0 in this subplot). (B) The modularity (averaged across densities) of the model networks at different values of  $\theta$ , which superimposes noise on top of the predicted relationships. For  $\theta = 0.18$ , the modularity of the model networks intersects with that of the actual  $\rho_q$ network. (C) Probability density plots of the average within-module correlation in gene expression (r<sub>GE</sub>, see Fig. 3): for the actual  $\rho_q$  network modules (black line); for modules derived from 1000 randomly generated networks (light gray distribution); and for modules derived from 1000 modeled networks with the same modularity as the actual  $\rho_q$  network (dark gray distribution).

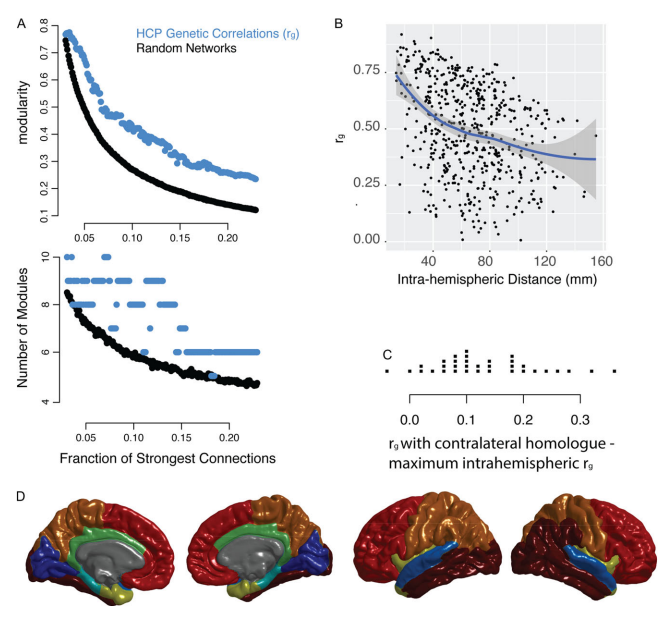


Figure 5. Replication of key results using data from Human Connectome Project. (A) Network models (graphs) were constructed at intervals of 0.001 connections density, where the connections density refers to the fraction of the strongest connections that are included in the graph. The genetic correlation networks are shown in addition to random control networks (see Methods). The top plot shows modularity, a measure of the strength of the community structure of the network, which is significantly higher in the genetic correlation network compared to the random network. The bottom plot shows the optimal number of modules at each density. For comparison with GOBS data, see Figure 1B. (B) An illustration of the consensus partition (modular community structure) with 8 cortical modules (For comparison with GOBS data, see Fig. 1C). (C) The relationship between anatomical distance and the genetic correlation between brain regions, shown for the left hemisphere. The line represents smoothed local averages, with standard error silhouettes (ggplot2 package in R). The right-hand plot focuses on the genetic correlation ( $\rho_a$ ), showing the fit of a quadratic model of the relationship with distance, which is a better fit than either a linear model or an exponential decay model (see results). For comparison with GOBS data, see Figure 2A. (D) For each of the FreeSurfer regions, the strip chart shows the difference between  $\rho_a$  with the contralateral homolog and the maximum within-hemisphere  $\rho_a$ .

significantly lower than for the  $r_{GE}$  within the actual  $\rho_a$  modules (P = 0.001, 1000 random networks). In contrast, for models with network modularity the same as the actual  $ho_q$  network, the within-module  $r_{GE}$  was similar to the within-module  $r_{GE}$  for the actual  $\rho_q$  modules (P = 0.4, 1000 network models).

#### Discussion

Network analysis of genetic correlations in human cortical thickness reveals a multiscale modular architecture. Different modular organizations, with greater or smaller numbers of regions per module, exist at different scales (or resolutions). In contrast to previous reports (Chen et al. 2012, 2013), we do not find evidence for a more "natural" number of clusters or modules at a certain spatial resolution. While perhaps not as intuitive as a single, unique modular structure, multiscale modularity has been reported by several brain functional and anatomical studies (Betzel and Bassett 2016). This structure may point to a hierarchical organization where larger modules are at least partially composed of smaller modules across spatial scales (Simon 1962).

Two fundamental principles appear to underlie this multiscale modular structure. First, regions are strongly genetically correlated

with their counterparts in the opposite cerebral hemisphere (contralateral homology; Levin 2005). Notably, heterogeneity in this contralateral homology, with lower homology in inferior prefrontal language areas, suggests a connection between functional and genetic asymmetry. Second, regions are highly genetically correlated with nearby regions, with an initial steep decrease with anatomical distance followed by a more gradual decline. These findings are replicated in an independent dataset (Glasser et al. 2013) and largely convergent with postmortem correlations in gene expression (Hawrylycz et al. 2012). Computational models demonstrate that, together, the constraints of contralateral homology and anatomical distance predict the degree of modularity of the actual genetic correlation network.

Physically embedded networks like the brain tend to have spatial constraints, with increased connectivity at shorter connection distance (Bullmore and Sporns 2012). Both brain functional (Alexander-Bloch et al. 2013) and diffusion imaging (Gong et al. 2012) networks are spatially constrained, which acts to decrease network wiring cost, although long distance connections do exist within functional modules (such as anterior and posterior regions of the default mode network). A possible molecular model of spatial constraints in genetic correlations is suggested by animal studies, with signaling molecules emitted by patterning centers in the developing cortex in a gradated fashion, and subsequently refined into disjunctive cortical maps based on sensory, thalamocortical connections (O'Leary et al. 2007). It should be noted that confounds such as motion artifact (Blumenthal et al. 2002; Power et al. 2012) and spatial smoothing by image processing pipelines could augment this signal, although this kind of artifact should not have differential impact on genetic as opposed to environmental correlations, and the regions we report with the strongest spatial constraints do not converge with those predicted to be differentially impacted by motion artifact in terms of cortical thickness estimates (Reuter et al. 2015; Alexander-Bloch et al. 2016).

Modules discovered by community detection techniques, which split heterogeneous data into relatively homogenous subgroups, are not necessarily significant in a statistical or biological sense (Lancichinetti et al. 2008; Porter et al. 2009). Robustness across a range of methods, and correspondence with known functional-anatomical boundaries, are some indicators of high-quality modular partitions. Evidence for a particular resolution of the community structure may be compelling (a specific, preferential number of modules), but multiscale systems by definition lack a specific resolution or scale. An advantage of network analysis is that the degree of modularity can be quantified relative to a null model. In the present case, for example, modularity is significantly higher than in random networks but not significantly higher than networks with the same anatomical distance constraints as the actual network, suggesting that the relationship with anatomical distance could account for the network's modularity.

Arguably this discussion begs the question of what counts as a "modular" network. Early work on modularity suggested standardized cut-offs for network modularity, e.g., Q > 0.3; however, because networks generated at random can have elevated modularity, current work stresses the importance of comparing the modularity of a network to that of a null model (Guimerà et al. 2004). While we interpret the present results as suggestive that anatomical constraints may account for the modularity of the genetic correlation network, a reasonable alternative conclusion is that relative to the computational null model (which preserves the spatial constraints and the

constraints of contralateral symmetry of the real data), the genetic correlation networks were not found to be modular.

The comparisons between neuroimaging data and gene expression analysis provides a potential link to biological mechanisms. For example, recent studies have shown an association between imaging-derived longitudinal changes in myelination and expression of genes related to synaptic activity in the same regions (Whitaker et al. 2016), as well as increased correlations in gene expression within resting-state fMRI networks for a subset of genes preferentially expressed in human upper cortical layers (Krienen et al. 2016). The fact that correlations in gene expression show analogous spatial constraints and relationships with contralateral homologs, compared to patterns of genetic correlation, also suggests that these relationships are not merely an artifact of imaging or postprocessing techniques. Similarly, the fact that modules derived from the imaging-genetics data show differential expression of genes related to nervous-system and telencephalon development is suggestive of an underlying biological mechanism. However, gene ontological analysis may be strongly influenced by publication bias and should be interpreted as hypothesisgenerating rather than confirmatory analyses. Correspondence between genetic cortical thickness modules and other ways of dividing the cortex into functionally or anatomically relevant subsections, such as the recent multimodal parcellation of the HCP data (Glasser et al. 2016), suggests the possibility of similar genetic principles of organization across modalities. Although beyond the scope of this paper, this hypothesis should be tested in future work.

Several additional and related methodological issues should be highlighted. (1) We do not directly assess the question of brain parcellation, i.e., how best to divide the brain into distinct regions. Although similar methods can be used to address this question, it is a mistake to conflate community detection with parcellation, as modules can be composed of spatially discontiguous regions. In addition, the degree of spatial smoothing necessary to make intersubject comparisons in morphology at the millimeter scale can bias inter-regional comparisons. (2) While the majority of published work on genetic correlations has used twin designs, analyses of extended pedigrees are superior in many ways including increased power and ability to distinguish genetic and environmental effects (Blangero 2004), as shared environment is necessarily reduced in extended pedigrees relative to twin pairs. The fact that the HCP data (unlike GOBS) is largely made up of twin pairs and siblings raises particular questions about the importance of distinguishing shared environment from genetic effects in these data. Because the GOBS data comes from extended pedigrees (Olvera et al. 2011), they do not share the same level of environmental overlap as sibling or twin samples, which makes replication in both datasets particularly compelling. Empirically, the fact that the genetic correlations are not notably higher in the HCP data than in the GOBS data (Figs. 1-2 and Fig. 5) also argues against the possibility of inflated estimates of genetic correlation due to these additional shared environmental factors. This is consistent with prior imaging-genetics work reporting similar genetic effects in "AE" structural equation models (which model additive genetic and individual environmental effects) compared to "ACE" models (which also model shared environmental effects), as estimates of common environmental variance tended to be at or near zero (Chen et al. 2012, 2013).

Another limitation (3) relates to the precision of the genetic correlations. The precision of an estimate of genetic correlation depends on the heritabilities of both traits, such that it is

estimated poorly when the traits have low heritabilities. Although mean regional cortical thicknesses of all regions were significantly heritable, there was some variability in their heritability values, and consequently, some variability in the precisions of their estimated genetic correlations. (This concerns also applies to prior published work on genetic correlations.) Finally, (4) there is no single correct way to account for both negative and positive connections in brain networks. Depending on the context, it may be appropriate to use network measures that can be estimated with both positive and negative edges (Traag and Bruggeman 2009), to include only positive edges, or to take the absolute value of edges (for example to focus on the magnitude of the shared genetic effect rather than the direction of the effect). In the present case, the results were robust to this methodological choice as the negative correlations were all lower in magnitude than the positive correlations which were included in the networks used to derive genetic modules.

In conclusion, it is worth stressing that the analysis of the genetic architecture of cortical thickness has potentially large clinical implications. Determined by the thickness of specific horizontal layers, which differ between regions in terms neuronal populations as well as neuropil (Rakic 1988), cortical thickness decreases in normal adolescence, in neurodegenerative disease (where loss of thickness is thought to represent neuronal death) and in schizophrenia (where loss of thickness may correspond to loss of neuropil in accordance with the overpruning hypothesis). Decreased thickness in schizophrenia, in particular, has been suggested to target specific "developmental modules," which develop in synchrony during normative adolescence (Alexander-Bloch et al. 2014) and are likely composed of regions influenced by overlapping sets of genes. In addition, schizophrenia is associated with alterations in the normative spatial constraints on anatomical (Bassett et al. 2008) and function connectivity (Alexander-Bloch et al. 2013), as well as disruption in normative constraints on the contralateral symmetry of morphology between hemispheres (Crow et al. 1989; Oertel-Knöchel and Linden 2011). Neuropsychiatric risk is thus likely to be expressed in over- or under-realization of normative organizational principles.

# **Supplementary Material**

Supplementary data is available at Cerebral Cortex online.

# **Notes**

Conflict of interest: None declared.

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