

## ORIGINAL ARTICLE

# Differential Contribution of Cortical Thickness, Surface Area, and Gyrification to Fluid and Crystallized Intelligence

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## Abstract

Human intelligence can be broadly subdivided into fluid (*gf*) and crystallized (*gc*) intelligence, each tapping into distinct cognitive abilities. Although neuroanatomical correlates of intelligence have been previously studied, differential contribution of cortical morphologies to *gf* and *gc* has not been fully delineated. Here, we tried to disentangle the contribution of cortical thickness, cortical surface area, and cortical gyrification to *gf* and *gc* in a large sample of healthy young subjects ( $n = 740$ , Human Connectome Project) with high-resolution MRIs, followed by replication in a separate data set with distinct cognitive measures indexing *gf* and *gc*. We found that while gyrification in distributed cortical regions had positive association with both *gf* and *gc*, surface area and thickness showed more regional associations. Specifically, higher performance in *gf* was associated with cortical expansion in regions related to working memory, attention, and visuo-spatial processing, while *gc* was associated with thinner cortex as well as higher cortical surface area in language-related networks. We discuss the results in a framework where “horizontal” cortical expansion enables higher resource allocation, computational capacity, and functional specificity relevant to *gf* and *gc*, while lower cortical thickness possibly reflects cortical pruning facilitating “vertical” intracolumnar efficiency in knowledge-based tasks relevant mostly to *gc*.

**Key words:** cognitive capacity, cortical gyrification, cortical surface area, cortical thickness, human intelligence

## Introduction

General intelligence (*g*) is frequently decomposed into fluid (*gf*) and crystallized (*gc*) components (Cattell 1963). While *gf* is associated with problem solving independent of previous knowledge, *gc* reflects acquired knowledge and general facts (Deary and Caryl 1997). Although they are correlated, they tap into distinct cognitive abilities and show different trajectory with aging, with *gf* generally declining with aging, while *gc* tending to increase with education and age (Horn and Cattell 1967; Manard

et al. 2014). Abilities related to *gf*, such as working memory, attention, and mental manipulation, are generally thought to benefit from a higher number of processing units, allowing for a more flexible allocation of resources during hypothesis testing (Unsworth et al. 2014; Duncan et al. 2017). On the contrary, knowledge-based cognitive performance relies on life-long learning, which is associated with optimization of existing computational schemes, where increased efficiency might be as important as computational power (Wenger et al. 2017a).

Identifying the neuroanatomical correlates of human intelligence has a long history. The initial attempts to relate brain anatomical measurements with intelligence started in the 19th century by relating head size to intelligence (Pearson 1906). These studies found a modest correlation between head size and intelligence, accounting for about 4% of variance (Pietschnig et al. 2015). The advent of neuroimaging techniques made it possible to measure brain volume noninvasively and relate regional cortical gray matter volume to intelligence measures, leading to the formalization of the well-known parieto-frontal integration theory (Jung and Haier 2007). While such evidence has helped localizing pivotal regions supporting *g* in humans, the exact nature of the contribution provided by specific structural properties of the brain is still unclear. Previous investigations on the association of cortical morphologies and human intelligence have mainly focused on gray matter volume and cortical thickness and have found inconsistent results across studies (Choi et al. 2008; Karama et al. 2011, 2014; Schnack et al. 2015).

Recent studies have shown that cortical gray matter volume can be decomposed into cortical thickness and cortical surface area, which are more biologically plausible morphological measures of cortical architecture (Winkler et al. 2010). Cortical surface area and cortical thickness have distinct developmental trajectories and uncorrelated genetic backgrounds (Panizzon et al. 2009). Cortical thickness is associated with radial neuronal migration and the number of neurons, dendritic arborizations, and glial support in cortical columns. In contrast, cortical surface area is related to tangential neuronal migration and captures the number of mini-columnar units in the cortex (Chenn and Walsh 2003; Rakic 2009; Rakic et al. 2009). Parsing the effect of cortical thickness and surface area on cortical volume has also functional relevance. For instance, cortical surface area and cortical thickness in human early visual cortices have been shown to exert opposite influences on neural population tuning with behavioral consequence for perceptual discrimination (Song et al. 2015). Therefore, in exploring the neural substrate of *gf* and *gc*, assessing the contribution of surface area and thickness seems important.

Aside from thickness and surface area, cortical folding, or gyrification, has also been proposed as a candidate neuroanatomical correlate of intelligence. Comparative anatomical studies have shown positive correlation between overall cortical gyrification and cognitive abilities across species (Zilles et al. 2013). Only recently, the regional association between gyrification and intelligence has been carved out, showing higher gyrification in large portions of parietal and frontal regions in subjects with higher *g* score (Gregory et al. 2016).

Human Positron Emission Tomography (PET) and task-based functional magnetic resonance imaging (fMRI) studies have shown activation of lateral prefrontal as well as parietal regions while subjects perform intelligence tasks. More recently, advances in human resting-state fMRI studies have shown the importance of specific functional networks including the frontoparietal network (FPN), default mode network (DMN), and dorsal attention network (DAN) in explaining individual variability in human intelligence (Colom et al. 2010; Santarnecchi et al. 2017a). The extent to which structural correlates of intelligence correspond to functional correlates of intelligence could offer new insight into structure–function relationship in the neuroscience of human intelligence.

In this study, we tested the hypothesis that *gf* and *gc* load on distinct cortical morphologies with different patterns of regional associations. To test this hypothesis, we first used a large sam-

ple of young healthy subjects from the Human Connectome Project (HCP,  $n = 740$ ) (Van Essen et al. 2013) with high-resolution structural T1-weighted MRIs and cognitive tests (identification phase). MRIs were analyzed using FreeSurfer pipeline (Fischl 2012), and a vertex-wise analysis followed by cluster-wise correction was pursued to identify the most significant clusters for each cortical morphology (cortical surface area, cortical thickness, and cortical gyrification). Afterwards, we showed that the structural correlates of intelligence follow the topography of resting-state human brain networks implicated in fluid and crystallized intelligence. Finally, we sought to replicate the findings of the identification phase in a separate data set (NKI-Rockland data set) (Nooner et al. 2012) with different cognitive tests in order to identify the most generalizable clusters linked to *gf* and *gc*.

## Materials and Methods

### Identification Data Set

A total of 740 subjects (age range, 21–35; right-handedness, >40) from the HCP who had 3T structural T1-weighted MRIs as well as neuropsychological data were included for this study.

### Behavioral Measures

The standard test for measuring fluid intelligence is the Raven's Standard Progressive Matrices (RSPM), a 60-item multiple-choice test with increasingly difficult pattern-matching tasks. The test assesses abstract reasoning with little dependency on language abilities. In the HCP, *gf* was measured using an abbreviated version of RSPM called Penn Progressive Matrices (PMAT). PMAT consists of 9-item subset of RSPM, which predicts the original RSPM with great accuracy (Bilker et al. 2012). We used the number of correct responses in the PMAT test for this study (PMAT24\_A\_CR, ranging from 0 to 9). The PMAT24\_A\_CR test has also been previously used in functional imaging studies (Finn et al. 2015). Also, a composite *gf* score (CompGfScore) derived from National Institutes of Health (NIH) toolbox (including the Dimensional Change Card Sort Test, the Flanker Inhibitory Control and Attention Test, the Picture Sequence Memory Test, the List Sorting Working Memory Test, and the Pattern Comparison Processing Speed Test) was tested (Akshoomoff et al. 2013), which gave similar results to PMAT24\_A\_CR. Here, we only report results for PMAT24\_A\_CR, which is considered a more direct measure of *gf* task.

*Gc* was measured using the NIH toolbox composite scores (Crystallized Cognition Composite), which combines Picture Vocabulary Test and Oral Reading Recognition Test into one score (CompGcScore) (Akshoomoff et al. 2013). The Oral Reading Recognition Test consists of reading and pronouncing either English or Spanish words as accurately as possible. The Picture Vocabulary Test measures receptive vocabulary using a computerized test by asking the subject to identify the correct picture for the audio recording of a word (for detailed description of the HCP behavioral measures, see <https://wiki.humanconnectome.org>).

### MRI Data Acquisition

Structural T1-MRIs in HCP were acquired using a dedicated customized 3-T Connectome Skyra scanner with the following parameters: MPRAGE sequence, TR = 2400 ms, TE = 2.14 ms, and



TI = 1000 ms; flip angle = 8 degrees; FOV = 224 × 224 mm; and voxel size = 0.7 mm isotropic.

## Image Processing and Statistical Analyses

Structural T1-weighted MRIs were processed with FreeSurfer v6.0 software package to create three-dimensional representations of cortical surface. FreeSurfer pipeline includes automated Talairach transformation, segmentation of subcortical white matter and deep gray matter structures based on intensity and neighbor constraints, intensity normalization, tessellation of gray matter–white matter boundary and gray matter–cerebrospinal fluid boundary, automated topology correction, and reconstruction of cortical surface meshes (Dale et al. 1999). Afterwards, reconstructed white matter surfaces were registered to FreeSurfer template (fsaverage) based on cortical folding patterns using spherical registration implemented in FreeSurfer (mri\_surf2surf). Cortical morphologies were then smoothed using a 10 mm full-width-at-half-maximum Gaussian kernel for thickness and surface area and 5 mm for gyrification. Cortical thickness was calculated at each point (vertex) in the cortex by measuring the distance between white and pial surfaces. The vertex-wise cortical surface area was calculated by averaging the area of all faces that meet at a given vertex on the white matter surface. The vertex-wise cortical gyrification was measured by calculating the gyrification index in circular three-dimensional regions of interest (Gregory et al. 2016). This method uses an outer smooth surface tightly wrapping the pial surface and computes the ratio between areas of circular regions on the outer surface and their corresponding circular patches on the pial surface. To make sure that our results are replicable using a different version of FreeSurfer, we also used FreeSurfer processed files provided by the HCP (db.humanconnectome.org) using FreeSurfer v5.3-HCP. In HCP, both T1-weighted and T2-weighted MRIs were used for cortical surface reconstruction. T2-weighted MRIs are mainly used to refine pial surface reconstruction.

In order to find the regional association of cortical morphologies with intelligence measures, a vertex-wise analysis followed by cluster-wise correction was pursued in the identification data set. We devised a method that enables better delineation of most significant vertices in cluster-wise correction and enables more stable cluster visualizations across different cortical morphologies (cortical thickness, surface area, and gyrification). We first split the data into 10-folds (each fold, 740/10, 74 subjects). While holding 1-fold out, we fit a linear regression model to all the remaining folds (9-folds, 666 subjects) at each vertex to test the association between vertex cortical morphometric and *gc/gf*, controlling for the effects of age, sex, and total brain volume (TBV). The results were then cluster-wise corrected (vertex-wise threshold, 0.01; cluster-wise *P* value, 0.05). We repeated this step 10 times, each time holding 1-fold out. We then concatenated all the resultant clusters across all iterations. The final map has values between 0 and 10 at each vertex, with 0 indicating that the vertex has not been selected in any of the iterations and 10 indicating that the vertex has been present in all the tested iterations. To show the usefulness of our approach, we also applied vertex-wise thresholding followed by cluster-wise correction on the whole sample (*n* = 740) and compared the final clusters. To make sure that our findings are not due to chosen vertex-wise threshold, we also performed our analyses

using vertex-wise *P* value of 0.001. The clusters obtained from applying both thresholds are provided as supplementary materials.

Although the effect size is considered small, TBV has been one of the most reliable and consistent predictor of intelligence. Given that it is a matter of controversy whether TBV should be included as a cofactor or not, we ran a follow-up analysis without controlling for TBV. Finally, performance in *gc* task is correlated with education level by the nature of the tasks, but education is not expected to affect *gf*. We first checked whether there is an association between education and *gf* (PMAT) and ran the analyses only on subjects who had at least 13 years of education. Finally, as both *gf* and *gc* are correlated with each other, we performed the analyses for *gf* controlling for *gc* and vice versa to find the most specific clusters associated with *gf* and *gc*.

## Overlap with Resting-State fMRI Networks

We tested the overlap between putative resting-state fMRI human networks and regional cortical morphology associated with *gf* and *gc*. We used 7 resting-state human networks previously derived from group-level analysis of 1000 subjects (Yeo et al. 2011). Dice coefficient (DC) was used to measure the overlap, defined as:

$$DC = 2 * N_{\text{overlap}} / (N_{\text{network}} + N_{\text{morphology}}).$$

$N_{\text{network}}$  is the number of vertices in the tested network,  $N_{\text{morphology}}$  is the number of significant vertices in the tested cortical morphology, and  $N_{\text{overlap}}$  is the number of vertices in common between the tested network and cortical morphology.

## Replication Data Set

To test the generalizability of the structural correlates, we tried to replicate the HCP clusters in the INDI data set, which offer distinct neuroimaging and behavioral measurements. High-quality structural T1-weighted MRIs (3T) for 253 subjects (age range, 21–45) were acquired using a 3-T Siemens MAGNETOM TrioTim with the following parameters: MPRAGE sequence, TR = 1900 ms, TE = 2.52 ms, and TI = 900 ms; flip angle = 9 degrees; FOV = 250 × 250 mm; voxel size = 1 mm isotropic.

Structural T1-MRIs were processed similar to HCP data. *gf* and *gc* were measured using the Wechsler Abbreviated Scale of Intelligence, a cognitive battery composed by 4 subtests: vocabulary, similarities, block design, and matrix reasoning. Vocabulary and similarities are combined to index *gc*, while block design and matrix reasoning are combined to form *gf*. Similar to identification phase, we first ran a vertex-wise analysis controlling for the effect of age, sex, and TBV. Afterwards, clusters present in at least 7 iterations (arbitrary threshold) from identification phase were tested separately for their corresponding cognitive measures, while controlling for the effects of age, sex, and TBV.

## Results

### Global Neuroanatomical Correlates of Intelligence

The total cortical surface area and average gyrification showed positive correlations with *gf* and *gc*, while average cortical

thickness was not correlated with *gf* or *gc* (Fig. 1). Given that global neuroanatomical measures might obscure the underlying associations, further testing of regional association of each cortical morphology with *gf* and *gc* was conducted.

### Structural Correlates of *Gf* and *Gc*

Figure 2 shows regional morphological associations with *gf* and *gc*. Cortical surface area in left and right superior parietal, left supramarginal, left caudal middle frontal, left pars-opercularis,

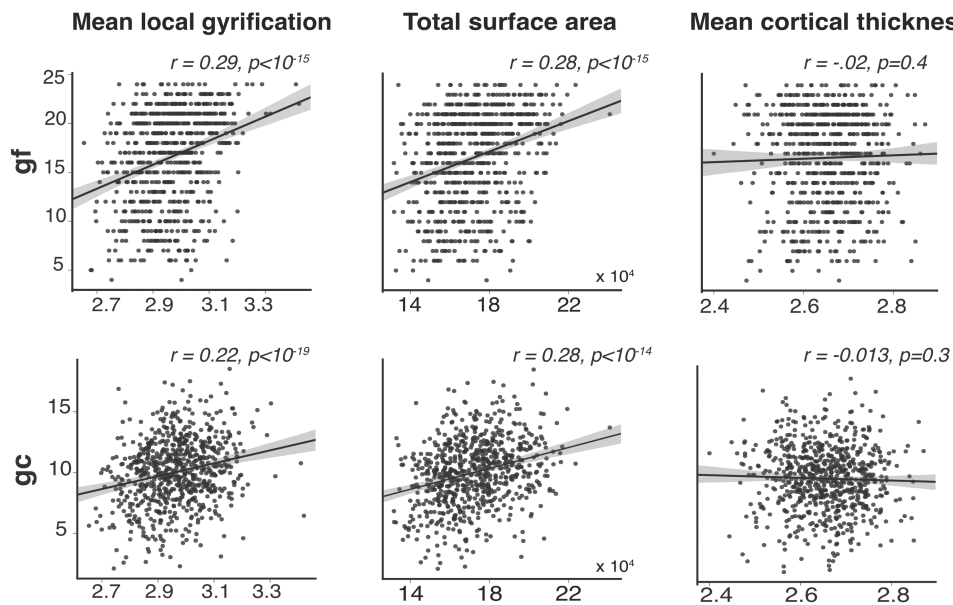


Figure 1. Global neuroanatomical correlates of *gf* and *gc*. Total surface area and mean gyrification are positively correlated with *gf* and *gc*. In contrast, mean cortical thickness showed no statistically significant association with *gf* or *gc*.

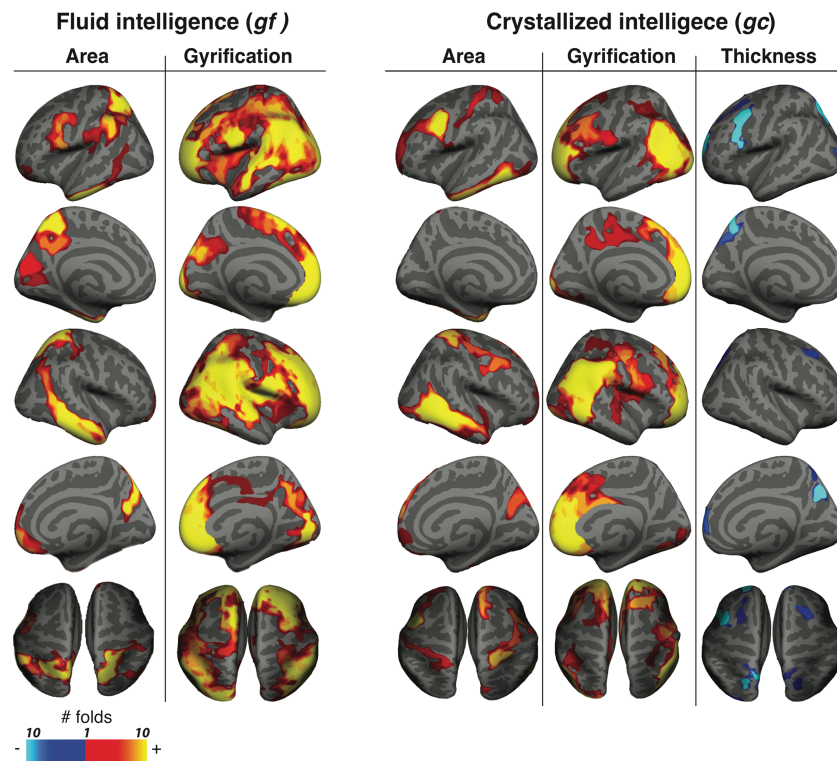


Figure 2. Regional morphological correlates of *gf* and *gc*. Vertex-wise structural correlates of *gf* and *gc* for cortical surface area, gyrification, and thickness. The subjects were divided into 10-folds. Vertex-wise analysis followed by cluster-wise correction was applied to combined subjects from 9-folds, while holding 1-fold out. The procedure was repeated 10 times. The resulting clusters were combined across all the iterations. The color shows the number of iterations that the vertex was within a statistically significant cluster (minimum, 0; maximum, 10). The red-yellow vertices show positive correlation; blue-cyan vertices show negative correlation.

left inferior temporal, right inferior and middle temporal, right medial orbitofrontal, and right rostral middle frontal were all positively correlated with *gf*. Cortical gyrification in large portions of left and right frontal, left and right inferior parietal, supramarginal, superior parietal, and superior temporal were positively correlated with *gf*. Cortical thickness did not show any regional association with *gf*. [Supplementary Figure S1](#) shows the results for the *gf*-associated cortical morphologies controlled for *gc*.

Cortical thickness in left caudal middle frontal, left pars-opercularis, left rostral middle frontal, left superior parietal, and right precuneus were all negatively correlated with *gc*. Cortical surface area in left and right caudal middle frontal, left inferior temporal, right middle temporal and right postcentral, and right superior frontal were all positively correlated with *gc*. [Supplementary Figure S1](#) shows the results for the *gc*-associated cortical morphologies controlled for *gf*.

Excluding TBV as a cofactor in the model did not influence the association between *gf/gc* and cortical thickness, while the association between *gf/gc* and cortical surface area and gyrification became statistically more significant with larger clusters ([Supplementary Fig. S2](#)).

We found a positive correlation between *gf* score indexed by PMAT24\_A\_CR and years of education ([Supplementary Fig. S3](#)). We ran the analyses only on subjects who had at least 13 years of education, to minimize the possible confounding effect of

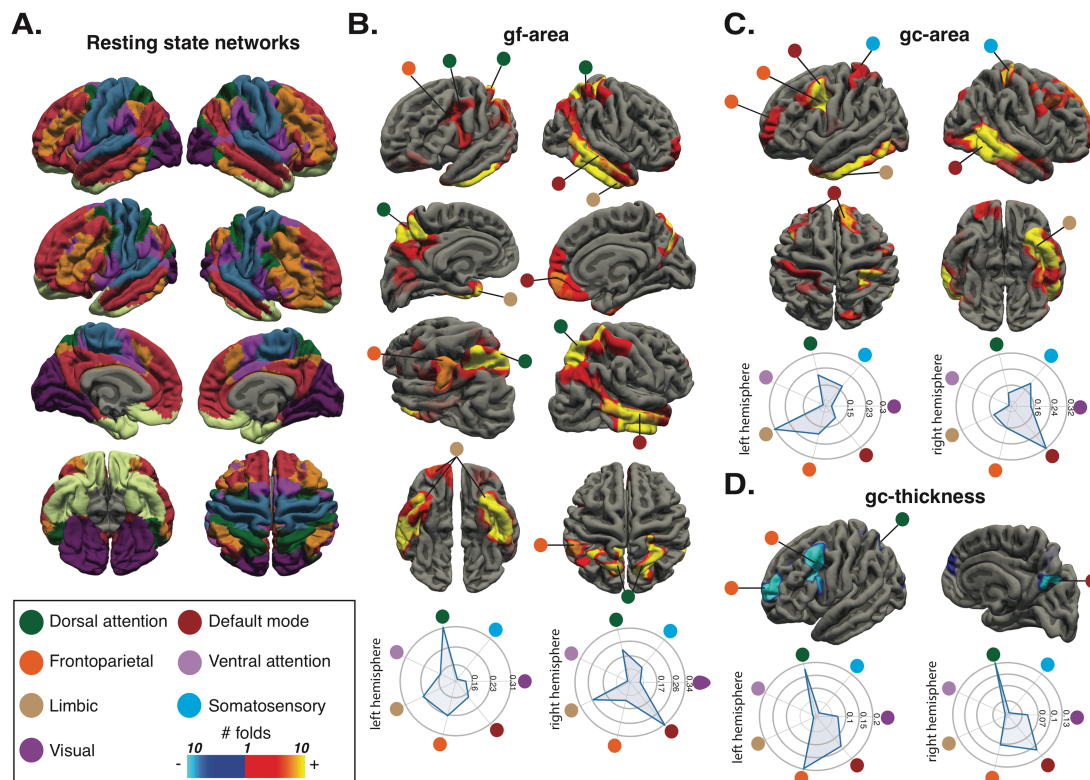
education level. The resulting clusters were similar to our original findings ([Supplementary Fig. S4](#)).

Clusters obtained from applying vertex-wise thresholding and cluster-wise correction to the whole data set ( $n = 740$ ) were similar to identical to clusters obtained from splitting the data into 10-folds ([Supplementary Fig. S5](#)). However, by splitting the data, we were able to also map the most important vertices within the clusters, providing final weighted cluster maps ([Supplementary Fig. S6](#)).

We also performed the analyses using a more stringent vertex-wise threshold of 0.001, with essentially similar results. We have provided the final clusters from applying both vertex-wise thresholds ( $P = 0.01$  and  $P = 0.001$ ) and the corresponding cluster-wise  $P$  value in a supplementary excel file. As a final sanity check, we redid the entire analyses using cortical surfaces reconstructions provided by the HCP and found similar to identical results ([Supplementary Fig. S7](#)).

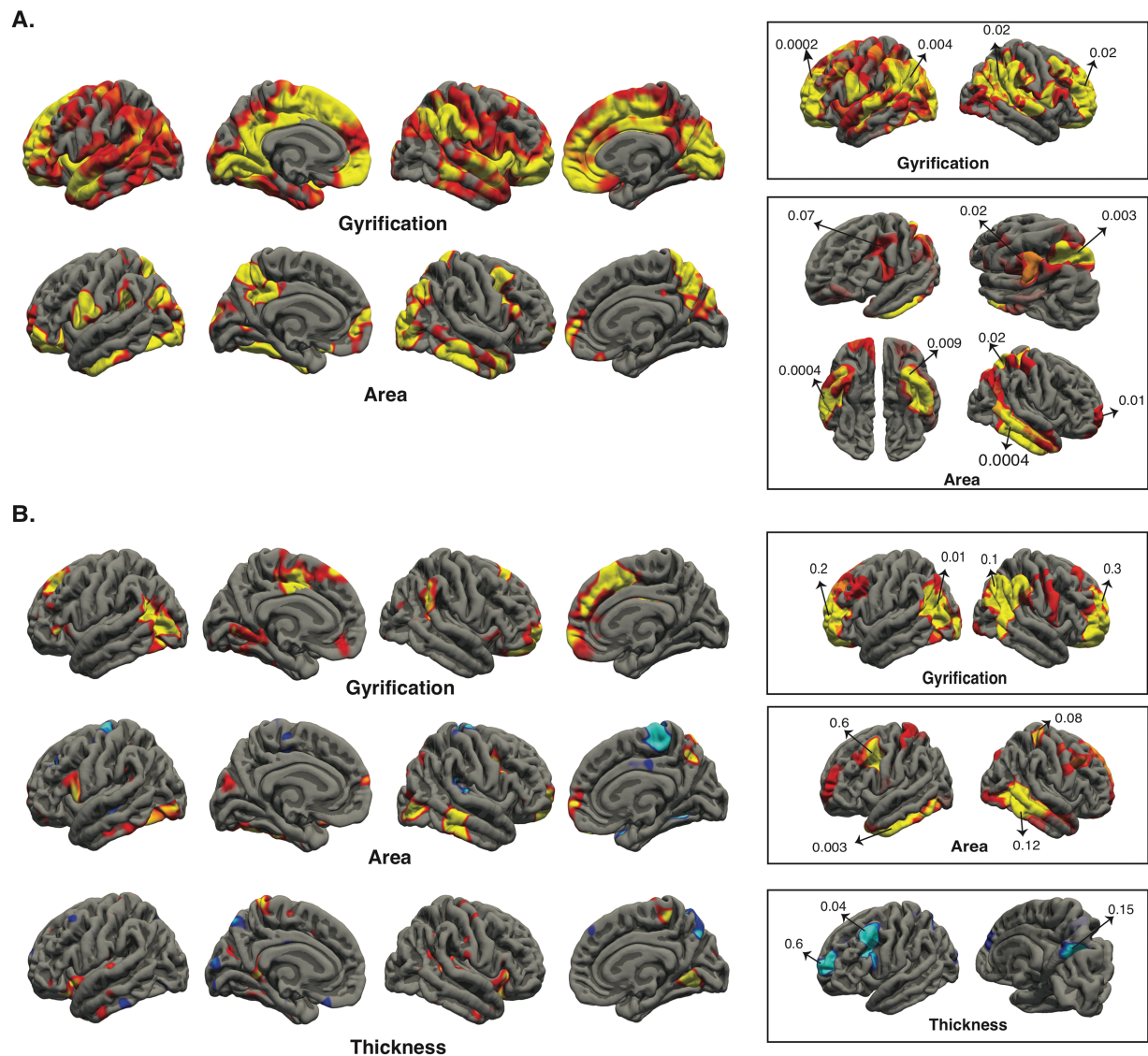
### Overlap with Resting-State fMRI Networks

We measured the overlap between resting-state fMRI networks and the regional cortical morphology associations of *gf* and *gc* ([Fig. 3](#)). *Gf*-related areas had the largest overlap with the DAN ( $DC = 0.31$ ) in the left hemisphere and the DMN ( $DC = 0.34$ ) in the right hemisphere. *Gc*-related areas had the largest overlap with the limbic ( $DC = 0.30$ ) in the left hemisphere and the DMN ( $DC = 0.32$ ) in the right hemisphere. *Gc*-related thickness had the



**Figure 3.** Correspondence between structural correlates of intelligence and functional resting-state human networks. Group-average ( $N = 1000$ ) resting-state human networks by [Yeo et al. \(2011\)](#) (A). *gf*-cortical surface area associations (B), *gc*-cortical surface area (C), and *gc*-cortical thickness associations (D). The radar graphs show the DC for each network. The circles show the correspondent network.





**Figure 4.** Replication of *gf* and *gc* clusters in INDI data set. Vertex-wise analysis ( $P \leq 0.05$ ) followed by testing the most significant clusters of identification phase in INDI for *gf* (A) and *gc* (B). The boxes show the tested HCP clusters in INDI, and the numbers are P values.

largest overlap with the FPN ( $DC=0.19$ ) in the left hemisphere and the DAN in the right hemisphere ( $DC=0.17$ ).

### Replication in INDI

Vertex-wise analyses for surface area, thickness, and gyrification were performed for *gf* and *gc* in INDI data set. Afterwards, significant clusters from identification phase (HCP) were separately tested in INDI data set for their corresponding cognitive measures. Similar to the identification phase, *gf* showed strong positive association with surface area in superior parietal, inferior temporal, and inferior frontal gyrus; positive association with gyrification in distributed regions of the cortical ribbon; no association with cortical thickness. Almost all tested clusters from the identification phase had statistically significant association with *gf* in INDI (Fig. 4A). Unlike *gf*, *gc* showed less striking association with cortical morphologies, and few clusters

from the identification phase showed statistically significant association with *gc* (Fig. 4B).

### Discussion

In this study, we tried to parse the link between different cortical morphologies and individual variability in *gf* and *gc* in a large data set of young healthy individuals, with replication in a separate data set using different behavioral measures. We found that while cortical gyrification in distributed regions of parietal, temporal, and frontal lobes displayed a positive correlation with *gf* and *gc*, cortical surface area and thickness showed more specific associations, which corresponded with meaningful cortical regions and resting-state fMRI networks. Specifically, *gf* showed positive association with cortical surface area and no association with cortical thickness. On the other hand, better performance in *gc* was associated with overall decreased cortical



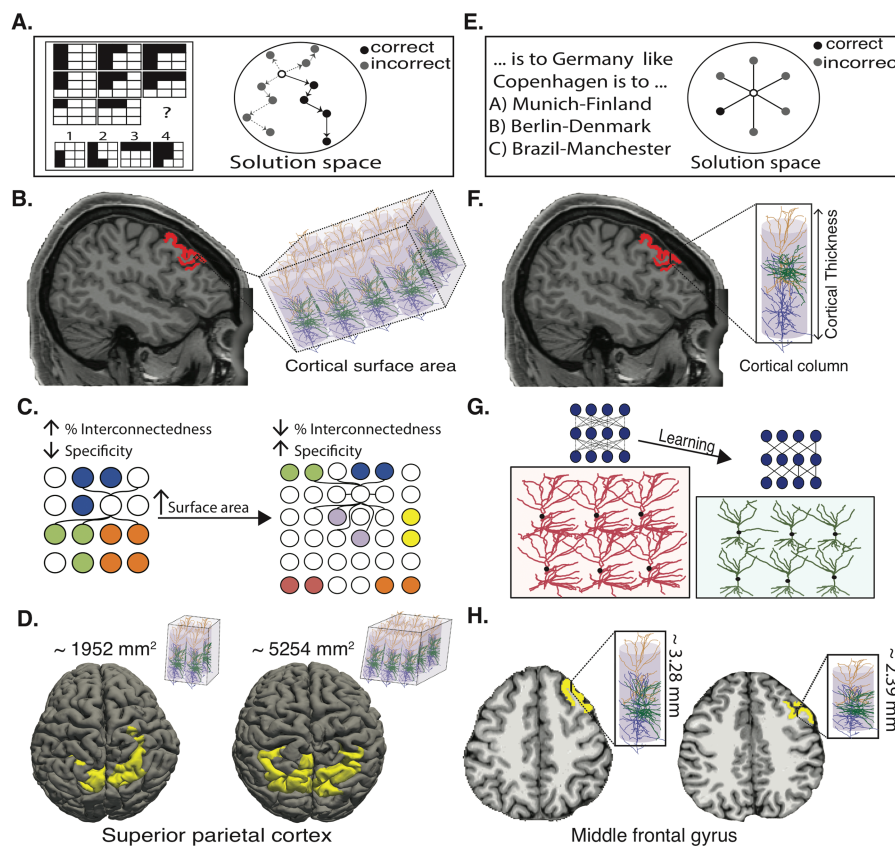
thickness as well as higher cortical surface area. Finally, higher surface area in left inferior temporal and right middle temporal gyri showed positive correlations with both *gf* and *gc*, suggesting possible shared structures contributing to both *gf* and *gc*.

### Computational Power versus Optimized Processing

*Gf* measures the ability to solve novel problems, which relies on generating, testing, and refuting multiple hypotheses to find the optimal solution to a given task (Fig. 5A). Cognitive factors underlying *gf* have been matters of study for decades. Importantly, performance in *gf* is highly correlated with working memory, suggesting that working memory capacity might be an important determinant of *gf* performance. The link between working memory performance and *gf* has been attributed to attention control, capacity to maintain goals, subgoals and hypotheses, and long-term memory (Unsworth et al. 2014). Moreover, functional imaging studies have shown that working memory, attention, and *gf* engage similar networks, suggesting possible shared neural correlates (Gray et al. 2003; Santarnecchi et al. 2017a). In this study, we found that better performance in *gf* was associated with cortical expansion in brain regions previously attributed to *gf*, working memory, attention as well

as visuo-spatial processing in structural and functional imaging (Gray et al. 2003; Colom et al. 2007; Kravitz et al. 2011, 2011; Petersen and Posner 2012; Constantinidis and Klingberg 2016), cortical morphologies (Deary et al. 2010), and lesion studies (Woolgar et al. 2010; Barbey et al. 2014).

How can regional cortical expansion lead to better performance in *gf*? Our brain has finite capacity for information processing (Luck and Vogel 1997). While the exact nature of the limited capacity is still not understood, two main models have been proposed. One model claims that there are only a limited number of available slots for holding information, and once filled, new information cannot be added (fixed capacity model) (Fukuda et al. 2010). In contrast, the second model argues that although there is no upper limit for number of representations, the competition between representations for resource allocation leads to information loss as the number of representations grow (flexible resource model) (Franconeri et al. 2013). Recent human psychophysics studies and monkey's electrophysiology have found more evidence in favor of the flexible resource model (Buschman et al. 2011; van den Berg et al. 2012; Miller and Buschman 2015). Regardless of the exact mechanism of capacity limit, higher cortical surface area can lead to increased information processing capacity by providing



**Figure 5.** Contribution of cortical thickness and surface area to *gf* and *gc*. Schematic example of a *gf* task, which requires navigating the solution space to find the correct answer (A). Cortical surface area is associated with the number of cortical columns (B). Higher cortical surface area can potentially lead to increased capacity by increasing the number of cortical columns (information processing units) as well as to increased functional specificity of cortical columns by reducing fraction of intercolumnar interconnections (C). Cortical surface area in superior parietal cluster morphed back into pial surface for two representative subjects (D). Schematic example of a *gc* task, which relies on accessing stored knowledge (E). Cortical thickness is associated with thickness of cortical columns determined by dendritic arborizations and glial support (F). Learning can lead to pruning of random connections between neurons, creating a more structured neural network, leading to decreased cortical thickness by reducing dendritic arborizations (G). Cortical thickness in left middle frontal gyrus morphed back into pial surface for two representative subjects (H).

more cortical columns (Rakic 2008), which are the functional units of the cortex (Mountcastle 1997) (Fig. 5B). Moreover, higher number of cortical columns should be associated with reduced fraction of intercolumnar connections (percent interconnectedness) (Ringo 1991) (Fig. 5C). This has been suggested to increase functional specificity of cortical columns and reduction of overlap in representations, which confer higher capacity to hold information (Ringo 1991). This idea has been recently supported by functional imaging study of neural population tuning in human early visual cortex: higher cortical surface area in early visual cortex is positively associated with sharpness of neural population tuning and better performance in perceptual discrimination (Song et al. 2015). By the same token, cortical expansion in *gf*-related brain regions can lead to higher information processing capacity and ability to generate (represent), test and refute multiple hypotheses simultaneously, with less interference between generated hypotheses (Franconeri et al. 2013). By parsing cortical volume into thickness and surface area, we were able to show that cortical expansion is the underlying factor for better performance in *gf* rather than higher cortical thickness (Fig. 5D). These results indicate the importance of studying cortical surface area as a separate entity in future *gf* studies.

Unlike *gf*, *gc* is related to accessing stored knowledge and general facts, which have been accumulated throughout years of learning and education (Deary et al. 2010) (Fig. 5E). In this study, we found that better performance in *gc* was associated with lower cortical thickness and higher cortical surface area. Previous studies on cortical changes associated with human learning and plasticity have shown increased cortical thickness (and volume, which mainly is driven by change in thickness) shortly after learning. Recently, the expansion-renormalization model has been hypothesized to explain unfeasible endless expansion of cortex (increased cortical thickness) in life-long learning (Wenger et al. 2017a). According to this model, learning-induced increased cortical thickness, brought by many cellular mechanisms including formation of new synapses and dendritic arborizations as well as proliferation of astrocytes, is followed by selection of the most efficient neural circuitries through pruning of inefficient connections (renormalization). The renormalization of cortical changes has been shown in human imaging studies (Wenger et al. 2017b) as well as animal models (Xu et al. 2009). Our findings support and extend the expansion-renormalization model. Repeated learning followed by pruning of unnecessary and weak connections between neurons would lead in long term to more stable and efficient neural circuitry, explaining decreased cortical thickness in the corresponding brain regions (Fig. 5G,H). Furthermore, our results tie in well with positive association of intelligence with both thinner cortex as well as faster rate of cortical pruning during development (Schnack et al. 2015) and less activation and metabolic consumption in individuals with higher cognitive performance (Haier et al. 1992).

Lower cortical thickness and higher cortical surface area in left caudal middle frontal gyrus were the most significant clusters for *gc*. Cortical thickness in left caudal middle frontal gyrus undergoes changes in cortical thickness in subjects learning new language (Mårtensson et al., 2012; Koyama et al. 2017). Higher cortical surface area accompanied by lower cortical thickness has been associated with better performance in visual perception discrimination and lend support to the proposed model of enlarged cortical surface area and decreased cortical thickness as the most advantageous cortical design for crystallized knowledge (Song et al. 2015).

## Shared Neuroanatomical Correlates of *gf* and *gc*

Although *gf* and *gc* rely on distinct cognitive abilities, individuals with higher *gf* perform better in *gc* tasks as well (Deary et al. 2010). This can imply engagement of some shared neural networks in both *gf* and *gc*. We specifically found that cortical surface area of anterior temporal lobe (including anterior inferior temporal gyrus) and middle temporal is associated with both *gf* and *gc*. Studies on patients with semantic dementia as well as human imaging studies have shown the role of anterior temporal lobe in formation of concepts and semantic knowledge (Ralph et al. 2017). Anterior temporal lobe specifically acts as a modality-invariant hub, which infers higher-order relationships among sensory, motor, verbal, and nonverbal information (Abel et al. 2015), and can play a significant role in both *gf* and *gc*.

Our results also show that cortical gyrification in frontal, parietal, and temporal regions was positively correlated with both *gf* and *gc*, which is in line with previous studies of regional cortical gyrification association with intelligence (Gregory et al. 2016). This indicates gyrification as a possible shared neuroanatomical correlate of both *gf* and *gc*, with less regional specifications but rather global characteristics of the brain.

In this study, we showed results with and without correction for TBV. Correcting for TBV reduced the size and significance level of the clusters for the association between *gf/gc* and cortical surface area. While cortical thickness shows more or less similar values across subjects with less amount of variability, cortical surface area varies greatly across subjects, which makes it a more prominent factor contributing to TBV. We believe that controlling for TBV more clearly reveals the regional association between cortical surface area and *gf/gc*, as the results are not contaminated by factors that are correlated with both TBV and intelligence, such as white matter volume.

## Brain Structure–Function Relationship and Human Intelligence

A central question in neuroscience is the relation between structure and function in the brain. We found that structural correlates of fluid and crystallized intelligence correspond to brain regions that are active while subjects perform fluid and crystallized intelligence tasks and to well-known human functional networks previously associated with human intelligence. Our results are also in line with the recent meta-analysis of *gf*-related co-activation maps showing great similarity with resting-state human brain networks including the DAN and the FPN (Santarnecchi et al. 2017b). These findings also provide structural evidence for the network-centric notion of human intelligence resulting from the interaction of multiple brain regions (Barbey 2018).

## Limitations of the Study and Future Directions

It is important to mention that metrics reported in this study are indirect measures of cortical morphologies using MRI. Importantly, it has been proposed that cortical thickness can be affected by myelination in deeper layers of the cortex especially during development (Sowell et al. 2001). Higher myelination in the gray matter–white matter boundary can lead to false segmentation of gray matter as white matter, resulting in apparent cortical thinning. Recently, thinning of visual cortex during development has been linked to increased myelination in the visual cortex, rather than pruning (Natu et al. 2018). Future studies using myelination maps combined with diffusion

imaging and T1-weighted MRI are needed to determine whether the negative association between cortical thickness and *gc* can be attributed to pruning or myelination.

Also, subject's movement during MRI acquisition can lead to underestimation of cortical thickness, which could be a potential confounding factor (Reuter et al. 2015). However, subjects with higher cognitive ability move less compared to subjects with lower cognitive ability (Wylie et al. 2014). This minimizes the possibility of movement as a confounding factor in our study, as the relationship between cortical thickness and *gc* was found to be negative.

Although we did not find any statistically significant association between cortical thickness and *gf*, this does not necessarily mean no association exists. Future studies with larger sample sizes are needed to further investigate the potential association between cortical thickness and *gf* with smaller effect sizes.

In this study, we were able to partially replicate results across two relatively large data sets, specifically for the association between *gf*, cortical surface area, and gyrification. However, results for the association between cortical morphologies and *gc* were not fully replicated. This can be partly due to relatively small sample size in the replication data set (INDI) and partly due to very different behavioral and psychological measurement of *gc* across two data sets. Future studies with larger sample size and more similar behavioral measures will reveal the generalizability of the findings across different data sets.

## Conclusion

We tried to disentangle the contribution of different cortical morphologies including cortical thickness, surface area, and gyrification, to individual variability in *gf* and *gc*. Our findings suggest a framework where “horizontal” cortical expansion, indexed by cortical surface area, enables greater resource allocation and computational capacity relevant to *gf* and *gc*, whereas decreased cortical thickness possibly reflects cortical pruning processes facilitating “vertical” intracolumnar efficiency in knowledge-based tasks linked to *gc*.

## Supplementary Material

Supplementary material is available at Cerebral Cortex online.

## Notes

The content of this paper is solely the responsibility of the authors and does not necessarily represent the official views of Harvard Catalyst, Harvard University and its affiliated academic health care centers, the National Institutes of Health and the Sidney R. Baer Jr. Foundation. *Conflict of Interest*: None declared.

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