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ORIGINAL ARTICLE

Fronto-Parietal Network Reconfiguration Supports the Development of Reasoning Ability

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Abstract

The goal of this fMRI study was to examine how well developmental improvements in reasoning ability can be explained by changes in functional connectivity between specific nodes in prefrontal and parietal cortices. To this end, we examined connectivity within the lateral fronto-parietal network (LFPN) and its relation to reasoning ability in 132 children and adolescents aged 6–18 years, 56 of whom were scanned twice over the course of 1.5 years. Developmental changes in strength of connections within the LFPN were most prominent in late childhood and early adolescence. Reasoning ability was related to functional connectivity between left rostrolateral prefrontal cortex (RLPFC) and inferior parietal lobule (IPL), but only among 12–18-year olds. For 9–11-year olds, reasoning ability was most strongly related to connectivity between left and right RLPFC; this relationship was mediated by working memory. For 6–8-year olds, significant relationships between connectivity and performance were not observed; in this group, processing speed was the primary mediator of improvement in reasoning ability. We conclude that different connections best support reasoning at different points in development and that RLPFC-IPL connectivity becomes an important predictor of reasoning during adolescence.

Key words: adolescent, child, development, dorsolateral, functional connectivity, inferior parietal lobule, parietal cortex, prefrontal cortex, processing speed, reasoning, rostrolateral, working memory

Introduction

Reasoning is among the most complex of cognitive operations that the human brain performs. It is also late-developing, with improvements in reasoning ability seen well into late adolescence (McArdle et al. 2002; Richland et al. 2006; Ferrer et al. 2009). Thus far, research on the neural underpinnings of reasoning ability has focused largely on age-related differences in the activation of specific brain regions (Wright et al. 2008; Crone et al. 2009; Eslinger et al. 2009; Dumontheil et al. 2010; Wendelken et al. 2011). However, the pattern of interaction between these regions is also critical, and recent investigations have begun to probe these interactions (Shokri-Kojori et al. 2012; Cocchi et al. 2013; Ebisch et al. 2013; Bazargani et al. 2014). Given the large improvements in reasoning ability that occur over childhood, a key question is whether and how these improvements are related to changes in functional connectivity-i.e., temporal correlation of activity between distinct brain regions. Here, we examine the relationship between functional connectivity and reasoning ability in a large longitudinal dataset, and investigate how changes in the pattern of network interactions over development support an improving capacity for reasoning.

Two broad regions, lateral prefrontal cortex (PFC) and posterior parietal cortex (PPC), are consistently reported in fMRI studies of reasoning (for review, see Prado et al. 2011; Krawczyk 2012; see Fig. 1). While several subregions within PFC and PPC are engaged during reasoning, we have argued that 2 regions in particular rostrolateral PFC (RLPFC) and the inferior parietal lobule (IPL) play a central role in reasoning about relations between stimuli (Wendelken et al. 2008, 2012; Wendelken and Bunge 2010; Watson and Chatterjee 2012; Vendetti and Bunge 2014). This conclusion is based on studies of matrix reasoning (Christoff et al. 2001; Kroger et al. 2002; Baldo et al. 2010), analogical reasoning (Bunge et al. 2005; Green et al. 2006; Wendelken et al. 2008; Cho et al. 2010; Volle et al. 2011), transitive inference (Wendelken and Bunge 2010; Waechter et al. 2013), and relational matching (Christoff and Gabrielli 2002; Bunge et al. 2009; Wendelken et al. 2012).

Age differences in the patterns of RLPFC and IPL activation have been reported in several prior fMRI studies involving child



Figure 1. The fronto-parietal network of regions commonly associated with reasoning, including approximate locations of left hemisphere ROIs examined in this study. Right hemisphere ROIs were a mirror image of these.

and adult participants (Wright et al. 2008; Crone et al. 2009; Dumontheil et al. 2010). In recent work, we demonstrated a shift in processing related to higher order relational reasoning, from predominantly dorsolateral prefrontal cortex (DLPFC) in younger children to predominantly RLPFC in older children and adolescents (Wendelken et al. 2011), similar to what we have observed previously in adults with the same task (Bunge et al. 2009). In parallel with this shift in lateral PFC, and increasing specialization of RLPFC for higher order relational reasoning, we observed increasing specialization of the IPL in the vicinity of the intraparietal sulcus, and a negative relationship between IPL cortical thickness and RLPFC functional selectivity.

To our knowledge, only one other study has examined age differences in functional connectivity in relation to reasoning ability. In an fMRI study of 37 older children and adults, Bazargani et al. (2014) observed a positive relationship between short-range (fronto-insular) connectivity and reasoning task performance, but no relationship between reasoning and long-range (frontoparietal) connectivity. Although the sample was not large enough to explore changing relationships across age groups, and the statistical analysis did not discriminate between specific region pairs, these results confirm the importance of inter-regional communication for reasoning performance.

While studies relating reasoning ability to functional connectivity are limited, a number of studies have examined the link between functional connectivity and IQ scores, which are based in part on measures of reasoning. One conclusion that emerges from this work is that better "global efficiency" of functional brain networks (i.e., a network with shorter paths for communication between nodes) is associated with better intellectual performance (Li et al. 2009; van den Heuvel et al. 2009; Langer et al. 2012). Consistent with this idea, but highlighting the role of one region in particular, Cole et al. (2012) demonstrated a relation between IQ and the "global brain connectivity" (i.e., average connectivity with the rest of the brain) of lateral PFC.

These functional connectivity studies suggest that improvements in reasoning ability might be best described by generalized increases in functional connectivity throughout the brain. In contrast, prior studies of relational reasoning in children and adults point toward a unique contribution of RLPFC and IPL. Thus, we

hypothesized that it is the development of this RLPFC-IPL functional connection that best supports age-related improvements in reasoning ability. Moreover, we hypothesized that there might be age-related shift in the specific connections whose strength best accounts for individual variability in reasoning ability. Specifically, we reasoned that as reasoning-related activation shifts forward from DLPFC to RLPFC, so too might the importance of DLPFC connectivity give way to rising importance of RLPFC connectivity. In the current study, we set out to test these hypotheses by examining the relationships between age, functional connectivity, and reasoning ability across ages 6–18. Further, given that the development of reasoning ability has been linked to improvements in processing speed and working memory (Kail and Salthouse 1994), we also included measures of these lower level cognitive abilities to determine whether any observed relations between functional connectivity and reasoning ability were mediated by processing speed and/or working memory.

Materials and Methods

Participants

This study included 132 typically developing individuals (76 males; 117 right-handed) from the Neurodevelopment of Reasoning Ability (NORA) study, a longitudinal project designed to examine the behavioral and neural factors that underlie changes in reasoning ability across childhood and adolescence. Analyses of the development of functional specialization and white matter microstructure based on the NORA study have previously been published (Wendelken et al. 2011; Ferrer et al. 2013). Data were available from all 132 participants at the initial assessment (T1) and from 56 of these participants at the second assessment (T2). At T1, participants ranged in age from 6 years to 18.7 years old $(11.1 \pm 3.6 \text{ years} \text{ [mean } \pm \text{ SD})$], while at T2, participants ranged in age from 7.3 to 19.1 $(11.9 \pm 3.2 \text{ years})$. The interval between assessments ranged between 0.9 and 2.2 years (1.5 \pm 0.3 years).

We tested for differences in attrition as a function of various measures, including age, gender, socioeconomic status (SES), and reasoning ability (see below), by regressing the number of time points on T1 values for each of these measures. There was no relationship between number of time points and age, gender, or reasoning score (all Ps > 0.2); however, there was a significant negative relationship between number of time points and SES (P = 0.015), indicating that lower SES participants were less likely than higher SES participants to return for T2 data collection.

Participants from the NORA project were included in the current study if they had usable fMRI data as well as data from 2 of 3 cognitive assessments of reasoning ability (at a given longitudinal time point). Scan data were considered usable if no more than 25% of total volumes in a given scan had volume-to-volume head motion in excess of 1 mm. All participants were screened for neurological impairment, psychiatric illness, history of learning disability, and developmental delay. All participants and their parents gave their informed assent (children under 12 years of age) or consent (adolescents aged 12 or more years and all parents) to participate in the study, which was approved by the Committee for Protection of Human Subjects at the University of California at Berkeley.

Behavioral Measures

Behavioral data were collected, at both T1 and T2, for a range of standardized cognitive tasks, including 3 standardized tests of reasoning ability: the "Block Design" and "Matrix Reasoning" subtests of the Wechsler Intelligence Scale for Children–Revised (WISC-R; Wechsler 1981), and the "Concept Formation" subtests of the Woodcock–Johnson Tests of Achievement (WJ-R; Woodcock and Johnson 1990). Block Design measures the ability to arrange a set of red-and-white blocks in such a way as to reproduce a 2D visual pattern shown on a set of cards. Matrix Reasoning measures the ability to select the geometric visual stimulus that accurately completes a series of stimuli that change along a particular dimension. Concept Formation measures the ability to identify and state the rules for concepts when shown illustrations of both instances and noninstances of the concept. All the tests are reported to have very high internal consistency and testretest reliability, ranging from 0.94 to 0.95 (McGrew et al. 1991; McArdle et al. 2002).

In addition to our measures of reasoning ability, we also examined processing speed and working memory. Our measure of processing speed was the Cross Out subtest of the WJ-R, which measures how accurately one can identify geometric shapes that match a sample stimulus. Our measure of working memory was the Digit Span task (from WISC-R), which involves remembering sequences of numbers in forward or backward order over a short delay.

MRI Data Acquisition

Brain imaging data were collected on a Siemens 3T Trio system at the UC Berkeley Brain Imaging Center. Functional imaging data analyzed here were collected in four 4-min scans, during which participants were engaged in a propositional visual analogy task, described below. Importantly, as discussed below, the data used in the present study was filtered so as to remove activations time-locked to the task. Behavioral and fMRI data associated with this task are the subject of a separate manuscript (Vendetti et al. under review).

The analogy task was identical in design to that of a prior study from our laboratory (Wright et al. 2008), with updated stimuli. Each run consisted of 10 semantic and 10 analogy trials, presented in random order in a fast, event-related design. Participants had up to 10 s to answer each question. On semantic trials, participants saw a target stimulus (e.g., a picture of a notepad) and selected which of 4 probe objects was most closely associated with it (e.g., a pen). On analogy trials, participants saw 3 target stimuli, arrayed as an incomplete analogy of the form A:B::C:?, and were asked to indicate which of 4 probe stimuli best completed the array. Age-related improvements were observed for both semantic and analogy trials; adult-level performance was reached at around age 10 for semantic trials, and around age 14 for analogy trials.

Scan parameters were as follows: gradient-echo EPI sequence, TR = 2000 ms, TE = 25 ms, 33 axial slices, $2.0 \times 1.8 \times 3.0$ mm voxels, no interslice gap, flip angle = 90°, field of view = 230 mm, 120 volumes per run). These 4 functional scans were preceded by collection of one or more high-resolution T₁-weighted MPRAGE anatomical scans (TR = 2300 ms, TE = 2.98 ms, $1 \times 1 \times 1$ mm voxels).

MRI Data Preprocessing

fMRI data preprocessing was carried out using SPM8 (Wellcome Trust Center for Neuroimaging, London). The first 3 volumes from each functional scanning run were discarded to allow for T1 equilibration. Functional images were corrected for differences in slice acquisition timing and were realigned to the first volume by means of 6-parameter rigid-body motion transformation. Motion parameters were extracted from this process and were used to inform a volume repair procedure (ArtRepair, Stanford Psychiatric Neuroimaging Laboratory). ArtRepair identified bad volumes on the basis of within-scan movement in excess of 1 mm and signal fluctuations in excess of 1.5 percent signal change, and then corrected bad volumes via interpolation. Following volume repair, the mean structural image was co-registered to the mean realigned functional image and then spatially normalized to SPM's T1 anatomical template. Normalization parameters obtained from this process were then applied to the un-normalized functional images to produce a set of functional images in SPM standard space (MNI152), with $3 \times 3 \times 3$ mm voxels. As a final preprocessing step, functional images were smoothed with an 8-mm FWHM istotropic Gaussian kernel.

Regions of Interest

To obtain an unbiased set of regions of interest (ROIs) that would accurately represent the regions within lateral PFC and PPC that we aimed to examine, we extracted coordinates from a large set of ROIs that have been used previously to examine global connectivity properties (Power et al. 2011). In particular, we selected coordinates that were representative of the rostral, dorsal, and ventral subdivisions of lateral PFC (RLPFC, DLPFC, and VLPFC, respectively) as well as coordinates associated with the inferior and superior parietal lobules (IPL and SPL, respectively) in the left hemisphere. To obtain comparable ROIs on the left and right sides, right hemisphere ROIs were constructed as mirror images of those on the left. MNI coordinates of each ROI are listed in Table 1, and approximate locations are shown in Figure 1. Each ROI was constructed as a sphere with 5 mm diameter. We measured the strength of functional connectivity for all pairs of ROIs within a hemisphere and for all homologous pairs across hemispheres.

Functional Connectivity Analysis

The purpose of the present study was to understand the relationship between reasoning ability and "intrinsic" patterns of functional connectivity—low-frequency correlations in regional activation that are relatively stable across task demands and that are thought to reflect the long-term history of coordination between regions (Seeley et al. 2007; Cole et al. 2014). Thus, we adopted the analytical methods of intrinsic functional connectivity analysis that are typically applied to resting-state data but that have also been successfully used with task data (e.g., Fair et al. 2007).

Intrinsic functional connectivity between regions was assessed by measuring low-frequency (f<0.08) correlations between BOLD activation time series extracted for each ROI, for

Table 1 Regions of Interest (ROIs) were constructed as 5-mm spheres centered at the given MNI coordinates, with right hemisphere ROIs at positive X values and left hemisphere ROIs at negative X values

ROI	Anatomical region	BA	Х	Y	Ζ
RLPFC	Anterior middle frontal gyrus	10	±34	55	4
DLPFC	Middle frontal gyrus	9	±42	25	30
VLPFC	Inferior frontal gyrus	45	±48	22	10
IPL	Inferior parietal lobe	40	±53	-49	43
SPL	Superior parietal lobe	7	±22	-65	48

BA, Brodmann's area. Coordinates are taken from Power et al. (2011).

each participant. Several additional preprocessing steps were undertaken to reduce spurious variance unlikely to reflect neural activity, including high-pass filtering (f > 0.009), regression of the global white matter signal averaged from a white matter ROI, regression of ventricular signal averaged from a ventricular ROI, and correction for motion effects (see below). Because we were working with fMRI task data (collected while participants performed an analogical reasoning task), we also regressed out task-related activity from each ROI time series (cf. Fair et al. 2007).

To address potential issues associated with head motion during fMRI data collection (Power et al. 2012; Satterwaite et al. 2012; Van Dijk et al. 2012), we employed 2 primary techniques. First, motion vectors, consisting of volume-to-volume movement in 6 dimensions (3 translation directions and 3 rotation axes), were regressed out of each ROI time series. Second, data scrubbing was implemented as a final preprocessing step (Power et al. 2012). This procedure deleted time points corresponding to bad volumes (as identified by ArtRepair) and yielded a concatenated time series for each ROI.

Statistical Analysis

To characterize reasoning ability in each participant, we first carried out a confirmatory factor analysis, using T1 data from the Matrix Reasoning, Block Design, and Concept Formation tasks, across the entire sample of participants. Factor loadings obtained from this analysis were combined with task scores from T1 and, separately, T2, in order to produce a reasoning ability factor score for each participant at each available time point. These factor scores were converted to z-scores for subsequent analyses. Factor loadings were similar for each task, so each contributed similarly to calculation of the reasoning ability factor score.

To examine age-related differences in reasoning ability, we fit a nonlinear model relating reasoning ability to age, utilizing the cumulative distribution function of the normal distribution ("pnorm" in R) as our nonlinear function. This function assumes that changes are monotonic with age, and also that, across the population, the age of maximal change is normally distributed around a population mean. There are 2 parameters in this model: age of maximal change (mean, μ) and spread (standard deviation, σ). To select optimal parameters for each modeled relation, we first considered only the T1 data and examined all possible values for μ between 6 and 19 years, in half-year increments, and all possible values for σ between half and 13 years, also in half-year increments. We then selected the parameters that yielded the maximum F-statistic and entered these optimal parameters into a nonlinear mixed model that included the full dataset (cross-sectional plus longitudinal), with age as a random factor conditioned on participant. These mixed-model analyses allowed us to account for within-individual changes while modeling age-related differences with all the available T1 and T2 data. Mixed-model analyses were conducted using the "nlme" package in R (R Development Core Team 2011).

To examine age-related differences in functional connectivity, we followed the same approach as with reasoning ability. First, we computed correlations between functional connectivity and age for each ROI pair, for the cross-sectional sample. Next, for each connection that demonstrated a significant correlation, we conducted a nonlinear mixed-model analysis with the full dataset.

To examine developing patterns of influence of connectivity on reasoning ability, we split our sample roughly evenly into 3 age groups based on age at T1, producing a younger third (N = 44, 6–8 years, mean \pm SD: 7.2 \pm 0.8 years), a middle third (N = 42, 9–11 years, 10.4 ± 0.9 years), and an older third (N = 46, 12–18 years, 15.4 ± 1.9 years). Specific age cutoffs were selected in order to roughly balance the group sizes. T2 data were available for 21 participants in the younger age group, 17 participants in the middle age group, and 18 participants in the older age group. Because participants were assigned to age groups on the basis of their age at T1, age ranges for each group were larger when T2 data were included: ages 6-10 for the younger group, 9-13 for the middle group, and 12-19 for the older group. For each age group, as well as for the full age range, and for each ROI pair, we conducted a multiple linear regression of reasoning ability on connectivity and age using T1 data, and a linear mixed-model regression of reasoning ability on connectivity and age using data from both longitudinal time points. Where effects were present, we included age group as indicator variable to test for an interaction between age group and connectivity in their effect on reasoning ability. In a follow-up analysis, in addition to examining connectivity of individual ROI pairs, we also computed average network connectivity across all ROI pairs and tested, as above, for effects of average network connectivity on reasoning ability.

To test whether or not processing speed or working memory mediate observed effects of functional connectivity on reasoning ability, we used structured equation modeling to relate age, functional connectivity, and reasoning ability with either processing speed or working memory. We limited our examination of this question to analysis of cross-sectional data and to those functional connections that demonstrated a relation with reasoning ability. For each connection, we used a χ^2 test to compare a full model that included a path from functional connectivity to reasoning ability with a reduced model that did not. The specific models are described in the Results section. Structured equation modeling was conducted with the "lavaan" package in R (Rosseel 2012).

Results

Development of Reasoning Ability

First, we examined age-related changes in reasoning ability. Fitting a nonlinear model to the cross-sectional data, we obtained an age of maximum change (μ) of 6 years with a spread (σ) of 4 years. Applying these parameters in a mixed-model regression to the full longitudinal dataset, we observed that this nonlinear relationship was highly significant (b = 5.3, t = 17.2, P < 0.001). Thus, these analyses indicate a rapid increase in reasoning ability during childhood, particularly before age 10, followed by a slower increase through adolescence (see Fig. 2).

Age-Related Differences in Functional Connectivity Within the Reasoning Network

We next sought to characterize patterns of developmental change in functional connectivity. A graphical summary of significant age-related changes is given in Figure 3A. Complete results, including cross-sectional correlations between age and connectivity as well as longitudinal mixed-model results, are reported in Table 2. In consideration of the 25 pairwise connections that were examined here, we consider findings with P < 0.002 as statistically significant, in accordance with the Bonferroni correction.

Within lateral PFC, a strong age-related increase in functional connectivity was observed between left DLPFC and left RLPFC (Fig. 3B), with best-fit parameters indicating maximal change at around age 13. In contrast, an age-related decrease in

connectivity was observed between DLPFC and VLPFC in both hemispheres. Using Fisher r-to-z transformation, we confirmed that the age-related patterns for DLPFC-RLPFC and DLPFC-VLPFC differed significantly from one another, on the left and on the right (Ps < 0.001). Connectivity between left and right DLPFC also demonstrated an overall increase with age. Within PPC, we observed a large decrease in functional connectivity between IPL and SPL in both hemispheres.



Figure 2. Age-related differences and longitudinal changes in reasoning ability, across the entire sample of participants. Age is on the x-axis, with reasoning ability factor score (RA) on the y-axis. Lines connecting data points indicate within-person longitudinal changes, while shape indicates the longitudinal time point (visit). The dotted line indicates the best-fitting cumulative normal distribution (with $\mu = 6$, $\sigma = 4$), and shading shows the standard error of this fit line.

A

Regarding fronto-parietal connectivity, we observed 2 notable age-related increases. For RLPFC, there was a strong age-related increase in connectivity with IPL, on the left (Fig. 3C) and on the right. This increase in IPL connectivity with RLPFC was significantly greater than changes in connectivity between IPL and either DLPFC (left: P = 0.006; right: P = 0.08) or VLPFC (left: P = 0.003; right: P = 0.02). For VLPFC, in contrast, there was an age-related increase in connectivity with SPL in both hemispheres. DLPFC demonstrated different patterns in the left and right hemispheres-specifically, increasing connectivity with SPL on the left and increasing connectivity with IPL on the right.

Examining the parameters of the best-fitting nonlinear models, we noted that age-related differences in functional connectivity were generally largest between the ages of 10 and 14, an age range that corresponds to pre- and early adolescence. This finding held for prefrontal-prefrontal differences (increases and decreases with age), for IPL-SPL age-related decreases, and for RLPFC-IPL age-related increases. There was greater variability in the timing of maximal change for other prefrontal-parietal connections, with right VLPFC-SPL increases peaking at around age 8.5 and right DLPFC-IPL increases peaking at around age 15.5.

Functional Connectivity and Reasoning Ability

Our primary aim was to understand how changing patterns of functional connectivity support developmental improvements in reasoning. We consider here the relation between reasoning ability and functional connectivity of different ROI pairs, across the entire age range and within each of the 3 age groups. Analyses are presented in the following order, according to our prior hypotheses: 1) fronto-parietal connections involving RLPFC, 2) fronto-frontal connections involving RLPFC, 3) connections



Figure 3. (A) Age-related differences in functional connectivity within the reasoning network. Regions are connected here if they demonstrated a significant pattern of agerelated changes (calculated using nonlinear mixed modeling). Thin lines indicate P < 0.05 uncorrected for multiple comparisons, thick lines indicate P < 0.05 Bonferronicorrected. (B) Scatter plot of the relationship between age left DLPFC-RLPFC connectivity. (C) Scatter plot of the relationship between age and left RLPFC-IPL connectivity.

Fable 2 Age-related	l differences	in functional	connectivity
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ROIs	Cross-sectional correlation			Longitudinal nonlinear mixed model				
	r	t	Р	μ	σ	b	t	Р
Prefrontal-prefrontal								
RLPFC-RLPFC (Bi)	-0.11	-1.3	0.18					
RLPFC-DLPFC (L)	0.30	3.6	0.0004	13	1	0.18	5.2	< 0.0001
RLPFC-DLFCP (R)	0.17	2.0	0.05	19	1	0.25	1.0	0.32
RLPFC-VLPFC (L)	-0.04	-0.50	0.61					
RLPFC-VLPFC (R)	-0.11	-1.3	0.19					
DLPFC-DLPFC (Bi)	0.23	2.7	0.008	14.5	1.5	0.17	4.1	0.0001
DLPFC-VLPFC (L)†	-0.26	-3.1	0.002	11	2.5	-0.14	-3.4	< 0.0001
DLPFC-VLPFC (R)	-0.34	-4.1	< 0.0001	13.5	2.5	-0.25	-6.1	< 0.0001
VLPFC-VLPFC (Bi)	-0.03	-0.38	0.70					
Parietal–parietal								
IPL-IPL (Bi)	0.15	1.8	0.07					
IPL-SPL (L)	-0.47	-6.1	< 0.0001	11	2	-0.32	-7.8	< 0.0001
IPL-SPL (R)†	-0.38	-4.6	<0.0001	13.5	2.5	-0.30	-5.0	< 0.0001
SPL-SPL (Bi)	-0.08	-1.0	0.33					
Prefrontal-parietal								
RLPFC-IPL (L)	0.39	4.8	<0.0001	12	2	0.25	6.1	< 0.0001
RLPFC-IPL (R)	0.37	4.5	<0.0001	12.5	1	0.25	6.2	< 0.0001
RLPFC-SPL (L)	0.19	2.1	0.03	11.5	1	0.11	2.8	0.008
RLPFC-SPL (R)	0.06	0.74	0.45					
DLPFC-IPL (L)	0.09	1.1	0.29					
DLPFC-IPL (R)	0.21	2.5	0.01	15.5	1	0.15	3.3	0.002
DLPFC-SPL (L)	0.27	3.1	0.002	11.5	1	0.10	2.8	0.006
DLPFC-SPL (R)	0.02	0.21	0.83					
VLPFC-IPL (L)	0.06	0.71	0.48					
VLPFC-IPL (R)	0.13	1.5	0.12					
VLPFC-SPL (L)	0.27	3.1	0.002	12.5	1	0.15	4.6	< 0.0001
VLPFC-SPL (R)	0.25	3.0	0.003	8.5	1	0.18	4.1	< 0.0001

For the cross-sectional analysis, reported statistics include the correlation *r*-value, the t-value, and the associated *P*-value. For the nonlinear mixed-model analysis of longitudinal data, which used the cumulative distribution function of the normal distribution, 5 values are reported: μ , the best-fit age of maximal change; σ , the best-fit standard deviation or spread; *b*, the parameter estimate for the best-fitting model; the associated t-value; and the associated *P*-value. Reported *P*-values are uncorrected for multiple comparisons. Given the 25 comparisons reported here, we consider *P* < 0.002 to be significant, in accordance with the Bonferroni correction. Mixed-model analyses were performed only in cases where the cross-sectional correlation survived *P* < 0.05 uncorrected.

[†]Mixed model did not converge; reported values are from cross-sectional nonlinear regression.

involving DLPFC that are not included above. We report results from both cross-sectional regressions of reasoning ability on connectivity and age (denoted "cross") and corresponding longitudinal mixed-model regressions (denoted "long"), including parameter estimates (b), t-values, and uncorrected P-values. We report all effects with uncorrected P-values <0.1. We consider a result to be significant if it survives Bonferroni correction for multiple comparisons (across pairs of regions and age groups) as follows: 1) P < 0.004, for the 12 tests relevant to our primary hypothesis, involving RLPFC-parietal connectivity; 2) P < 0.003, for the 15 tests relevant to our secondary hypothesis, involving RLPFC-frontal connections; and 3) P < 0.002, for the 24 additional tests involving connections with DLPFC.

Based on prior findings demonstrating the involvement of left RLPFC and IPL in reasoning tasks, our main prediction was that reasoning ability would be supported by left RLPFC-IPL connectivity. Across the entire age range, this effect was not significant (cross: b = 0.47, t = 1.7, P = 0.09; long: P > 0.1). However, when this analysis was limited to 12–18-year olds, the effect of left RLPFC-IPL connectivity on reasoning ability was highly significant (cross: b = 1.7, t = 3.5, P = 0.001; long: b = 0.93, t = 3.5, P = 0.003; Fig. 4A). Moreover, there was an interaction between connectivity and membership in the older age group in their effect on reasoning ability (cross: t = 2.4, P = 0.02; long: t = 2.3, P = 0.02). The

correlation between left RLPFC-IPL connectivity and reasoning ability was not significant for either the younger or middle age groups, and the correlation between right RLPFC-IPL connectivity and reasoning ability was not significant for any age group (all Ps > 0.1).

Although we predicted that RLPFC-IPL connectivity would be most strongly related to reasoning ability, we also examined other fronto-parietal connections. Across the full age range, there was no effect of either left or right RLPFC-SPL connectivity on reasoning ability (all Ps > 0.1). But notably, for the older group, in stark contrast to the positive relationship between reasoning ability and left RLPFC-IPL connectivity, there was a weak negative relationship between reasoning ability and left RLPFC-SPL connectivity (cross: b = -1.0, t = -1.7, P = 0.08; long: b = -0.83, t = -2.5, P = 0.02; Fig. 4B). With regard to fronto-parietal connections involving DLPFC or VLPFC, we observed no significant relationship between connectivity and reasoning after accounting for age, either across the entire age range or within each age group, either cross-sectionally or longitudinally (all Ps > 0.1).

We turned next to consideration of RLPFC connectivity with other prefrontal regions. After accounting for age, there was no significant relationship between RLPFC-VLPFC or RLPFC-DLPFC connectivity and reasoning ability, left or right, across the entire age range or within any of the 3 age groups, cross-sectionally or



Figure 4. Graphs depicting the relationship between functional connectivity and reasoning ability (RA) in older children and adolescents. (A) A scatter plot of the positive relation between left RLPFC-IPL connectivity and reasoning ability. (B) A scatter plot of the negative relation between left RLPFC-SPL connectivity and reasoning ability.

longitudinally (all Ps > 0.1). However, there was a significant positive effect of cross-hemispheric connectivity between left and right RLPFC in the middle age group (cross: b = 1.1, t = 3.1, P = 0.003; long: b = 0.85, t = 3.0, P = 0.008). This effect was limited to the middle age group, as indicated by a significant interaction between group membership and connectivity (cross: t = 3.2, P = 0.002; long: t = 2.7, P = 0.009).

In contrast to RLPFC, cross-hemispheric connectivity was not associated with reasoning ability for DLPFC, VLPFC, IPL, or SPL (all Ps > 0.1). There were no other significant relationships between connectivity and reasoning ability, for any intraprefrontal, intraparietal, or fronto-parietal connections, for any age group (Ps > 0.07) [There was a weak negative relationship between left DLPFC-VLPFC connectivity and reasoning ability in the youngest group (cross: b = -0.68, t = -1.8, P = 0.07; long: did not converge). Although this was the strongest effect of connectivity on reasoning ability observed in the younger age group, it was not significant. This effect was not present in the other two age groups or across the entire age range (all Ps > 0.1)]. In addition, there was no effect of average network connectivity on reasoning ability in any group (all Ps > 0.1).

Mediation Analysis: Processing Speed and Working Memory

Given that the development of reasoning ability has been linked to the development of both processing speed and working memory (Fry and Hale 1996; Nettelback and Burns 2010), we sought to determine whether the observed relations between functional connectivity and reasoning ability were mediated by processing speed and/or working memory. We examined structured equation models that related reasoning ability, functional connectivity, and age, and which also included either processing speed or processing speed and working memory as potential mediators of the path from functional connectivity to reasoning ability. We compared each full model to a reduced model that eliminated the path from functional connectivity to reasoning ability. We report the χ^2 test for this model comparison along with key parameters from the full model here, and present more detailed results for the full models in Table 3. Summary results, depicting significant relationships for each group, are shown in Figure 5. We note that only a subset of participants completed the working memory task, so models involving working memory were tested with a reduced sample (numbers are shown in Table 3). It is for this reason that we considered processing speed separately, as well as in combination with working memory.

First we examined, for the older group, the connection between left RLPFC and left IPL that demonstrated a significant relation with reasoning ability. For the processing speed model, removing the path from functional connectivity to reasoning ability led to significant reduction in model quality ($\Delta \chi^2(1) = 10.9$, P < 0.001). Indeed, this path was highly significant in the full model (b = 0.28, P < 0.001). In contrast, the paths from processing speed to reasoning ability, and from functional connectivity to processing speed, were not significant (Ps > 0.2). Thus, processing speed was unrelated to either reasoning ability or RLPFC-IPL connectivity in the older group. For the working memory model, removing the path from connectivity to reasoning ability again led to a significant worsening of the model fit ($\Delta \chi^2(1) = 11.1$, P < 0.001), and this path in the full model was highly significant (b = 0.43, P < 0.001). The path from working memory to reasoning ability showed a negative trend (b = -0.12, P = 0.2), while the path from connectivity to working memory was marginal and positive (b = 0.46, P = 0.06). Thus, while working memory was weakly related to both reasoning ability and RLPFC-IPL connectivity in 12-18-year olds, it did not mediate the effect of connectivity on reasoning in this group (Fig. 5C).

Next, we examined the connection between left and right RLPFC that demonstrated a positive relation with reasoning ability in the middle age group. Here again, for the processing speed model, removal of the path from connectivity to reasoning ability resulted in a significantly worse model fit ($\Delta \chi^2(1) = 7.7$, P < 0.01). This path was highly significant in the full model (b = 0.25, P = 0.004), while the path from processing speed to reasoning ability was marginally significant (b = 0.25, P = 0.08). The path from connectivity to processing speed was not significant (b = 0.12, P = 0.18). Thus, the impact of bilateral RLPFC connectivity on reasoning ability in the middle age group is not mediated by processing speed. For the working memory model, the results were different. In this case, removal of the path from connectivity to

Table 3 Regression path param	eters associated with each or	otimal model from the me	diation analyses
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A PS model: RLPFC-IPL connectivity, ages 12–18 (N = 46) Full model RA ~ Age 0.08 0.18 0.45 0.66 RA ~ PS 0.20 0.19 1.04 0.30 RA ~ FC 0.28 0.08 3.52 <0.001 PS ~ Age 0.58 0.11 5.19 <0.001 PS ~ Age 0.35 0.06 0.74 0.46 FC ~ Age 0.35 0.26 1.43 0.15 S. PS and WM model: RLPFC-IPL connectivity, ages 12–18 (N = 20) Pull model RA Age 0.07 0.64 RA ~ PS 0.10 0.21 0.47 0.64 RA ~ PS 0.10 0.12 0.47 0.64 WM ~ PG 0.37 0.53 0.70 0.48 WM ~ PS 0.57 0.48 1.20 0.24 WM ~ PS 0.57 0.48 1.20 0.24 WM ~ PS 0.57 0.48 0.24 0.24 RA ~ Age 0.37 0.39 0.94 0.34 RA ~ PS 0.4 0.14 1.74 0.08 C. PS model	Regression path	Estimate	Std. Err.	Z-value	P-value
RA - Age 0.08 0.18 0.45 0.66 RA - PS 0.20 0.19 1.04 0.30 RA - PC 0.28 0.08 3.52 -0.001 PS - Age 0.58 0.11 5.19 -0.001 PS - FC 0.05 0.06 0.74 0.46 FC - Age 0.38 0.26 1.43 0.15 B. PS and WM model: RLPFC-IPL connectivity, ages 12-18 (N = 20) Full model RA -0.01 -1.28 0.001 RA - PS 0.10 0.21 0.47 0.64 0.20 RA -0.001 RA -0.23 .020 RA -PS 0.20 0.001 -1.28 0.20 RA -PS 0.21 0.01 -1.28 0.20 RA -PS 0.27 0.46 0.24 1.89 0.06 FC - Age 0.23 .0.70 0.48 0.24 0.23 .0.70 0.23 .0.70 0.23 .0.75 0.066 FC - Age 0.25 0.029 .1.18 0	A. PS model: RLPFC-IPL con	nectivity, ages 12–18 (N = 46) Full	model		
RA PS 0.20 0.19 1.04 0.30 RA PC 0.28 0.08 3.52 0.001 PS<	RA ~ Age	0.08	0.18	0.45	0.66
RA FC 0.28 0.08 3.52 0.001 PS - Age 0.05 0.06 0.74 0.001 PS - Age 0.38 0.26 1.43 0.05 PS - Age 0.38 0.26 1.43 0.01 PS - Age 0.76 0.23 3.32 0.001 RA - PS 0.10 0.21 0.47 0.64 RA - PC 0.43 0.11 3.86 -0.001 WM ~ Age 0.37 0.53 0.70 0.48 WM ~ Age 0.37 0.53 0.70 0.48 WM ~ PS 0.57 0.48 1.20 0.24 C - PS model: RLPFC-RLPFC connectivity, ages 9-11 (N = 42) Full model 74 0.44 0.24 C - PS model: RLPFC-RLPFC connectivity, ages 9-11 (N = 42) Full model 74 0.04 0.24 RA - PS 0.4 0.14 1.74 0.08 RA - PC 0.25 0.09 1.35 0.18 PS - Age 0.017 0.39 2.7	RA ~ PS	0.20	0.19	1.04	0.30
PS - Age 0.58 0.11 5.19 -0.001 PS - FC 0.05 0.06 0.74 0.45 E. PS and WM model: RLPFC-IPL connectivity, ages 12-18 (N = 20) Full model N 0.001 0.23 3.2 0.001 RA - Age 0.10 0.21 0.47 0.64 0.64 RA - VM -0.12 0.10 -1.28 0.001 WM - Age 0.37 0.53 0.70 0.48 WM - PS 0.57 0.48 1.20 0.24 WM - FC 0.46 0.24 1.89 0.06 FC - Age 0.37 0.53 0.70 0.48 RA - PS 0.4 0.14 1.74 0.89 RA - PS 0.4 0.49 0.41 0.44 0.24 RA - PS 0.4 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49 0.49	$RA \sim FC$	0.28	0.08	3.52	< 0.001
PS - FC 0.05 0.06 0.74 0.46 FC - Age 0.38 0.26 1.43 0.15 B. PS and WM model: RLPFC-IPL connectivity, ages 12-18 (N = 20) Full model 0.01 0.21 0.47 0.64 RA - Age 0.76 0.23 3.32 0.001 RA - PS 0.10 -1.28 0.20 RA - FC 0.43 0.11 3.86 -0.001 0.48 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.24 0.23 0.23 0.24 0.23 0.23 0.24 0.23 0.23 0.70 0.48 0.001 0.44 0.23 0.23 0.24 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.23 0.24 0.89 0.06 FC - Age -0.46 0.39 0.24 0.36 0.23 0.23 0.25 0.09 0.25 0.000 RA - FC 0.25 0.09 0.25 0.006 FC - Age 0.16	PS ~ Age	0.58	0.11	5.19	< 0.001
FC - Age 0.38 0.26 1.43 0.15 B. PS and WM model: RLPFC-IPL connectivity, ages 12-18 (N = 20) Full model 0.43 0.23 3.22 0.01 RA - Age 0.76 0.23 3.22 0.07 0.64 0.64 RA - NS 0.10 0.21 0.47 0.64 0.20 RA - FC 0.43 0.11 3.86 0.001 WM - Age 0.37 0.53 0.70 0.48 WM - PC 0.46 0.29 -1.18 0.24 C. PS model: RLPFC-RLPFC connectivity, ages 9-11 (N = 42) Full model RA - PS 0.46 0.39 -0.14 0.34 RA - PS 0.4 0.14 1.74 0.08 0.34 RA - PS 0.04 0.14 1.74 0.08 RA - PS 0.4 0.14 1.74 0.08 0.35 0.38 0.55 0.36 0.13 0.34 0.4 0.14 0.74 0.04 0.18 0.75 0.06 0.55 0.55 0.55 0.56 0.53 0.55 0.56 0.53 0.55 0.56 0.58 <	PS ~ FC	0.05	0.06	0.74	0.46
B. PS and WM model: RLPFC-IPL connectivity, ages 12-18 (N = 20) Full model 0.001 RA - Age 0.76 0.23 3.32 0.001 RA - Age 0.10 0.21 0.47 0.64 RA - FC 0.43 0.11 3.86 -0.001 WM - Age 0.37 0.53 0.70 0.48 WM - PS 0.57 0.48 1.20 0.23 WM - PS 0.57 0.48 1.20 0.23 WM - PS 0.57 0.48 1.20 0.23 WM - PS 0.57 0.48 1.20 0.24 C. PS model: RLPFC-RLPFC connectivity, ages 9-11 (N = 42) Full model 0.24 0.24 0.24 RA - Age 0.37 0.39 0.94 0.34 0.34 RA - Age 0.37 0.39 2.91 0.004 0.5 RA - Age 0.37 0.39 2.91 0.006 0.5 0.55 0.05 0.56 0.13 0.29 0.05 0.55 0.05 0.56 0.55 0.56 0.55 0.56 0.56 0.55 0.55 0.56	FC ~ Age	0.38	0.26	1.43	0.15
RA ~ Age 0.76 0.23 3.32 0.001 RA ~ PS 0.10 0.21 0.47 0.64 RA ~ PC 0.43 0.11 3.86 <0.001	B. PS and WM model: RLPFO	C-IPL connectivity, ages 12–18 (N	= 20) Full model		
RA~PS 0.10 0.21 0.47 0.64 RA~VM -0.12 0.10 -1.28 0.20 RA~FC 0.43 0.11 3.86 <0.001	RA ~ Age	0.76	0.23	3.32	0.001
RA~WM-0.120.10-1.280.20RA~FC0.430.113.86<0.001	RA ~ PS	0.10	0.21	0.47	0.64
RA~FC 0.43 0.11 3.86 <0.001 WM ~ Age 0.37 0.53 0.70 0.48 WM ~ PS 0.57 0.48 1.20 0.23 WM ~ PC 0.46 0.24 1.89 0.06 FC ~ Age -0.46 0.24 1.89 0.06 C. PS model: RLPFC connectivity, ages 9-11 (N = 42) Full model RA ~ Age 0.37 0.39 0.94 0.34 RA ~ PS 0.4 0.14 1.74 0.08 RA ~ PS 0.4 0.14 1.74 0.08 RA ~ FC 0.25 0.09 2.91 0.004 PS ~ Age 1.07 0.39 2.75 0.006 PS ~ Age 1.07 0.39 2.75 0.006 RA ~ Age 0.56 D.56 D.56 D.56 D.56 D.55 0.56 D.55 D.55 D.55 D.55 D.55	$RA \sim WM$	-0.12	0.10	-1.28	0.20
WM ~ Age 0.37 0.53 0.70 0.48 WM ~ PS 0.57 0.48 1.20 0.23 WM ~ PC 0.46 0.24 1.89 0.06 FC ~ Age -0.46 0.39 -1.18 0.24 C. PS model: RLPFC-RLPFC connectivity, ages 9-11 (N = 42) Full model 0.39 0.94 0.34 RA ~ Age 0.37 0.39 0.94 0.34 0.34 RA ~ Age 0.37 0.39 0.94 0.34 RA ~ PC 0.25 0.09 2.91 0.004 PS ~ Age 1.07 0.39 2.75 0.006 PS ~ Age 0.012 0.09 1.35 0.18 PC ~ Age -0.38 0.65 -0.58 0.56 D. PS and WM model: RLPFC-RLPFC connectivity, ages 9-11 (N = 31) Reduced model: RA ~ PC removed RA ~ Age 0.013 3.29 0.001 RA ~ Age -0.05 0.36 0.13 3.29 0.001 RA ~ PS 0.42 0.13 3.29 0.00	$RA \sim FC$	0.43	0.11	3.86	<0.001
WM ~PS 0.57 0.48 1.20 0.23 WM ~PC 0.46 0.24 1.89 0.06 FC ~Age -0.46 0.39 -1.18 0.24 C. PS model: RLPFC-RLPFC connectivity, ages 9-11 (N = 42) Full model 0.39 0.94 0.34 RA ~Age 0.37 0.39 0.94 0.34 0.34 RA ~PS 0.4 0.14 1.74 0.08 RA ~FC 0.25 0.09 2.91 0.004 PS ~Age 1.07 0.39 2.75 0.006 PS ~Age -0.38 0.65 -0.58 0.56 D. PS and WM model: RLPFC-RLPFC connectivity, ages 9-11 (N = 31) Reduced model: RA ~FC removed RA ~Age -0.03 0.02 RA ~Age -0.05 0.36 -0.13 0.90 RA ~Age -0.05 0.36 -0.13 0.90 RA ~PS 0.42 0.13 3.29 0.001 RA ~Age -0.05 0.36 -0.13 0.90 RA ~PS	$WM \sim Age$	0.37	0.53	0.70	0.48
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	WM ~ PS	0.57	0.48	1.20	0.23
FC~Age-0.460.39-1.180.24C. PS model: RLPFC-RLPFC connectivity, ages 9-11 (N = 42) Full model </td <td>WM ~ FC</td> <td>0.46</td> <td>0.24</td> <td>1.89</td> <td>0.06</td>	WM ~ FC	0.46	0.24	1.89	0.06
C. PS model: RLPFC-RLPFC connectivity, ages 9-11 (N = 42) Full model 0.39 0.94 0.34 RA ~ Age 0.37 0.39 0.94 0.34 RA ~ PS 0.4 0.14 1.74 0.08 RA ~ FC 0.25 0.09 2.91 0.004 PS ~ Age 1.07 0.39 2.75 0.006 PS ~ Age 0.12 0.09 1.35 0.18 FC ~ Age -0.38 0.65 -0.58 0.56 D. PS and WM model: RLPFC-RLPFC connectivity, ages 9-11 (N = 31) Reduced model: RA ~ FC removed 0.90 0.90 RA ~ Age -0.05 0.36 -0.13 0.90 RA ~ Age -0.05 0.36 -0.13 0.90 RA ~ PS 0.42 0.13 3.29 0.001 RA ~ WM 0.26 0.12 2.30 0.02 WM ~ Age 0.39 0.53 0.73 0.46 WM ~ Age 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.16 FS model: no connectivity, ages 6-8 (N = 49) Reduced model: RA ~ Age removed<	FC ~ Age	-0.46	0.39	-1.18	0.24
RA~Age 0.37 0.39 0.94 0.34 RA~PS 0.4 0.14 1.74 0.08 RA~FC 0.25 0.09 2.91 0.004 PS~Age 1.07 0.39 2.75 0.006 PS~Age 0.12 0.09 1.35 0.18 FC~Age -0.38 0.65 -0.58 0.56 D. PS and WM model: RLPFC-RLPFC connectivity, ages 9-11 (N = 31) Reduced model: RA~FC removed 0.90 RA~Age 0.90 RA~Age -0.05 0.36 -0.13 0.90 0.001 RA~Age -0.05 0.36 -0.13 0.90 0.001 RA~PS 0.42 0.13 3.29 0.001 RA~VM 0.26 0.12 2.30 0.02 WM~Age 0.39 0.53 0.73 0.46 WM ~PS -0.10 0.19 -0.56 0.58 WM ~PS -0.10 0.71 -1.41 0.16 F. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~Age removed RA ~PS 0.68 0.18 3.7 <0.001	C. PS model: RLPFC-RLPFC o	connectivity, ages 9–11 (N = 42) Fu	ll model		
RA~PS 0.4 0.14 1.74 0.08 RA~FC 0.25 0.09 2.91 0.004 PS~Age 1.07 0.39 2.75 0.006 PS~FC 0.12 0.09 1.35 0.18 FC~Age -0.38 0.65 -0.58 0.56 D. PS and WM model: RLPFC-RLPFC connectivity, ages 9-11 (N = 31) Reduced model: RA ~FC removed 0.90 RA ~PS 0.42 0.13 3.29 0.001 RA ~VM 0.26 0.12 2.30 0.02 0.02 WM ~Age 0.39 0.53 0.73 0.46 WM ~PS -0.10 0.19 -0.56 0.58 WM ~PS -0.10 0.71 -1.41 0.16 E. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~Age removed <0.001	RA ~ Age	0.37	0.39	0.94	0.34
RA~FC 0.25 0.09 2.91 0.004 PS~Age 1.07 0.39 2.75 0.006 PS~FC 0.12 0.09 1.35 0.18 FC~Age -0.38 0.65 -0.58 0.56 D. PS and WM model: RLPFC-RLPFC connectivity, ages 9-11 (N = 31) Reduced model: RA~FC removed 0.90 RA~Age -0.05 0.36 -0.13 0.90 RA~Age -0.05 0.36 -0.13 3.29 0.001 RA~Age 0.39 0.53 0.73 0.42 WM ~Age 0.39 0.53 0.73 0.46 WM ~PS -0.10 0.19 -0.56 0.58 WM ~FC 0.36 0.13 2.74 0.006 FC ~Age -1.0 0.71 -1.41 0.16 FS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~Age removed 3.7 <0.001	RA ~ PS	0.4	0.14	1.74	0.08
PS ~ Age 1.07 0.39 2.75 0.006 PS ~ FC 0.12 0.09 1.35 0.18 FC ~ Age -0.38 0.65 -0.58 0.56 D. PS and WM model: RLPFC-RLPFC connectivity, ages 9-11 (N = 31) Reduced model: RA ~ FC removed 0.90 RA ~ Age -0.05 0.36 -0.13 0.90 RA ~ PS 0.42 0.13 3.29 0.001 RA ~ WM 0.26 0.12 2.30 0.02 WM ~ Age 0.39 0.53 0.73 0.46 WM ~ PS -0.10 0.19 -0.56 0.58 WM ~ FC 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.66 F. PS model: no connectivity, ages 6-8 (N = 19) Reduced model: RA ~ Age removed <0.001	$RA \sim FC$	0.25	0.09	2.91	0.004
PS~FC 0.12 0.09 1.35 0.18 FC~Age -0.38 0.65 -0.58 0.56 D. PS and WM model: RLPFC-RLPFC connectivity, ages 9-11 (N=31) Reduced model: RA~FC removed 0.90 RA~Age -0.05 0.36 -0.13 0.90 RA~PS 0.42 0.13 3.29 0.001 RA~VM 0.26 0.12 2.30 0.02 WM~Age 0.39 0.53 0.73 0.46 WM ~PS -0.10 0.19 -0.56 0.58 WM ~PS -0.10 0.71 -1.41 0.16 FC ~Age -1.0 0.71 -1.41 0.16 FC ~Age -1.0 0.71 -1.41 0.16 FC ~Age -1.4 0.32 4.3 <0.001	PS ~ Age	1.07	0.39	2.75	0.006
FC~Age -0.38 0.65 -0.58 0.65 D. PS and WM model: RLPFC-RLPFC connectivity, ages 9-11 (N = 31) Reduced model: RA ~ FC removed 0.90 RA ~ Age -0.05 0.36 -0.13 0.90 RA ~ PS 0.42 0.13 3.29 0.001 RA ~ WM 0.26 0.12 2.30 0.02 WM ~ Age 0.39 0.53 0.73 0.46 WM ~ PS -0.10 0.19 -0.56 0.58 WM ~ FC 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.16 E. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~ Age removed <	PS ~ FC	0.12	0.09	1.35	0.18
D. PS and WM model: RLPFC-RLPFC connectivity, ages 9–11 (N = 31) Reduced model: RA ~ FC removed 0.00 RA ~ Age -0.05 0.36 -0.13 0.90 RA ~ PS 0.42 0.13 3.29 0.001 RA ~ WM 0.26 0.12 2.30 0.02 WM ~ Age 0.39 0.53 0.73 0.46 WM ~ PS -0.10 0.19 -0.56 0.58 WM ~ FC 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.16 F. PS model: no connectivity, ages 6–8 (N = 44) Reduced model: RA ~ Age removed 3.7 <0.001	FC ~ Age	-0.38	0.65	-0.58	0.56
RA ~ Age -0.05 0.36 -0.13 0.90 RA ~ PS 0.42 0.13 3.29 0.001 RA ~ WM 0.26 0.12 2.30 0.02 WM ~ Age 0.39 0.53 0.73 0.46 WM ~ PS -0.10 0.19 -0.56 0.58 WM ~ FC 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.16 E. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~ Age removed 3.7 <0.001	D. PS and WM model: RLPF	C-RLPFC connectivity, ages 9–11 (N = 31) Reduced model: RA ~ FC	removed	
RA~PS 0.42 0.13 3.29 0.001 RA~WM 0.26 0.12 2.30 0.02 WM ~ Age 0.39 0.53 0.73 0.46 WM ~ PS -0.10 0.19 -0.56 0.58 WM ~ FC 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.16 E. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~ Age removed RA ~ PS 0.68 0.18 3.7 <0.001	RA ~ Age	-0.05	0.36	-0.13	0.90
RA~WM 0.26 0.12 2.30 0.72 WM ~ Age 0.39 0.53 0.73 0.46 WM ~ PS -0.10 0.19 -0.56 0.58 WM ~ FC 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.16 E. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~ Age removed RA ~ PS 0.68 0.18 3.7 <0.001	RA ~ PS	0.42	0.13	3.29	0.001
WM ~ Age 0.39 0.53 0.73 0.46 WM ~ PS -0.10 0.19 -0.56 0.58 WM ~ FC 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.16 E. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~ Age removed RA ~ PS 0.68 0.18 3.7 <0.001	$RA \sim WM$	0.26	0.12	2.30	0.02
WM ~ PS -0.10 0.19 -0.56 0.58 WM ~ FC 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.16 E. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~ Age removed RA ~ PS 0.68 0.18 3.7 <0.001	$WM \sim Age$	0.39	0.53	0.73	0.46
WM ~ FC 0.36 0.13 2.74 0.006 FC ~ Age -1.0 0.71 -1.41 0.16 E. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~ Age removed RA ~ PS 0.68 0.18 3.7 <0.001	WM ~ PS	-0.10	0.19	-0.56	0.58
FC ~ Age -1.0 0.71 -1.41 0.16 E. PS model: no connectivity, ages 6-8 (N = 44) Reduced model: RA ~ Age removed <td>$WM \sim FC$</td> <td>0.36</td> <td>0.13</td> <td>2.74</td> <td>0.006</td>	$WM \sim FC$	0.36	0.13	2.74	0.006
E. PS model: no connectivity, ages 6–8 (N = 44) Reduced model: RA ~ Age removed 3.7 <0.001	FC ~ Age	-1.0	0.71	-1.41	0.16
RA ~ PS 0.68 0.18 3.7 <0.001 PS ~ Age 1.4 0.32 4.3 <0.001	E. PS model: no connectivit	y, ages 6–8 (N = 44) Reduced mode	el: RA ~ Age removed		
PS ~ Age 1.4 0.32 4.3 <0.001 F. PS and WM model: no connectivity, ages 6-8 (N = 19) Reduced model: RA ~ Age removed <td>$RA \sim PS$</td> <td>0.68</td> <td>0.18</td> <td>3.7</td> <td><0.001</td>	$RA \sim PS$	0.68	0.18	3.7	<0.001
F. PS and WM model: no connectivity, ages 6–8 (N = 19) Reduced model: RA ~ Age removed 0.23 0.02 RA ~ PS 0.61 0.26 2.38 0.02 RA ~ WM 0.19 0.16 1.22 0.22 WM ~ Age -0.36 0.99 -0.37 0.72 WM ~ PS 1.04 0.42 2.50 0.01 PS ~ Age 1.64 0.40 4.14 <0.001	PS ~ Age	1.4	0.32	4.3	<0.001
RA ~ PS 0.61 0.26 2.38 0.02 RA ~ WM 0.19 0.16 1.22 0.22 WM ~ Age -0.36 0.99 -0.37 0.72 WM ~ PS 1.04 0.42 2.50 0.01 PS ~ Age 1.64 0.40 4.14 <0.001	F. PS and WM model: no co	nnectivity, ages 6–8 (N = 19) Redu	ced model: RA ~ Age removed		
RA ~ WM 0.19 0.16 1.22 0.22 WM ~ Age -0.36 0.99 -0.37 0.72 WM ~ PS 1.04 0.42 2.50 0.01 PS ~ Age 1.64 0.40 4.14 <0.01	$RA \sim PS$	0.61	0.26	2.38	0.02
WM ~ Age -0.36 0.99 -0.37 0.72 WM ~ PS 1.04 0.42 2.50 0.01 PS ~ Age 1.64 0.40 4.14 <0.001	$RA \sim WM$	0.19	0.16	1.22	0.22
WM ~ PS 1.04 0.42 2.50 0.01 PS ~ Age 1.64 0.40 4.14 <0.001	$WM \sim Age$	-0.36	0.99	-0.37	0.72
PS~Age 1.64 0.40 4.14 <0.001	WM ~ PS	1.04	0.42	2.50	0.01
	$PS \sim Age$	1.64	0.40	4.14	<0.001

We tested for mediating effects of either processing speed (PS) alone, or processing speed and working memory (WM). (A and B) For older children and adolescents (ages 12–18), we examined mediating effects on the relationship between left RLPFC-IPL functional connectivity (FC) and reasoning ability (RA). (C and D) For the middle group (ages 9–11), we examined mediating effects on the relationship between bilateral RLPFC connectivity and reasoning ability. (E and F) For younger children (ages 6–8), we examined mediating effects on the relationship between age and reasoning ability. Only a subset of participants completed the working memory task; numbers of participants are listed for each model. Relationships between age, processing speed, and functional connectivity are not shown for the models that include working memory, as these relationships are likely to be more accurately estimated in the PS-only models.

reasoning ability did not significantly affect the model fit $(\Delta \chi^2(1) = 0.01, P > 0.2)$, and this path was not significant in the full model (b = 0.01, P > 0.2). However, the path from working memory to reasoning ability was significant (b = 0.26, P = 0.04), and there was a significant positive relationship between connectivity and working memory (b = 0.29, P = 0.008). Thus, working memory mediates the effects of bilateral RLPFC connectivity on reasoning ability in 9–11-year olds (Fig. 5B).

Finally, we sought to better understand the large age-related changes in reasoning ability in the youngest group, for whom changes in functional connectivity did not have a significant effect. Here, we sought to determine whether processing speed and/or working memory mediated the relationship between reasoning ability and age. We examined simple mediation models that included a direct path from age to reasoning ability as well as a mediating path through either processing speed or working memory, and compared these with reduced models that excluded the path from age to reasoning ability. In the processing speed model, the fit of the reduced model was marginally worse than that of the full model ($\Delta\chi^2(1) = 3.2$, P = 0.07). In the full model, the regression path from age to reasoning ability was similarly marginal (b = 0.82, P = 0.07) while the regression paths from age to processing speed (b = 1.4, P < 0.001) and from processing speed to reasoning ability (b = 0.68, P < 0.001) were highly significant. For the model that also included working memory, the reduced model was not significantly worse than the full model ($\Delta\chi^2(1) = 1.3$, P > 0.2). The regression path from age to reasoning ability in the full model was not significant



Figure 5. Summary of mediation analysis results, for models examining the relationships between age, functional connectivity, processing speed (PS), working memory (WM), and reasoning ability (RA). Dashed lines indicated relationships that do not hold in the presence of PS- and/or WM-mediating variables. (A) In younger children (6–8 years), processing speed mediated the relationship between age and reasoning ability. (B) In middle children (9–11), working memory mediated the relationship between functional connectivity (RLPFC bilateral connectivity) and reasoning ability. (C) In older children and adolescents (12–18 years), neither processing speed nor working memory mediated the relationship between functional connectivity (left RLPFC-IPL connectivity) and reasoning ability.

(P > 0.2). There was a significant path connecting processing speed to working memory (b = 1.04, P = 0.01), but only positive trends in the relationships between age and working memory (b = 1.0, P = 0.17) and between working memory and reasoning ability (b = 0.21, P = 0.18). Thus, in children 6–8 years old, processing speed was the primary mediator of developmental improvements in reasoning ability (Fig. 5A).

Discussion

Age-Related Differences in Functional Connectivity

We observed marked changes in the pattern of functional connectivity in the network of regions that were examined, from childhood through adolescence. Within lateral PFC, the predominant pattern was one of differentiation of DLPFC (and to a lesser extent RLPFC) from VLPFC, suggesting that the functions of DLPFC and VLPFC may become increasingly disparate over development. In contrast, the present results suggest increased communication between RLPFC and DLPFC. This emerging pattern is consistent with a hierarchical model of lateral PFC, in which RLPFC depends on outputs of DLPFC (Koechlin et al. 2004; Badre and D'Esposito 2007).

A second prominent pattern of observed age-related changes were the selective increases in fronto-parietal connectivity. In particular, the observed increase in RLPFC-IPL connectivity was expected, as IPL is the region that in many adult studies demonstrates the strongest functional connectivity with RLPFC (Vincent et al. 2008; Boorman et al. 2009). The increase in VLPFC-SPL connectivity observed here is not as clearly linked to previously reported connectivity patterns in adults. Overall, the developmental changes in fronto-parietal connectivity suggest increasing communication between prefrontal regions and specific parietal targets. The logical outcome of these changes, a system in which different parietal subregions, is consistent with reported parietal connectivity patterns in adults (Nelson et al. 2010; Mars et al. 2011).

Nonlinear modeling revealed that many of these age-related changes in connectivity peak between the ages of 10 and 14. This suggests that the transition to adolescence may be marked by widespread changes in patterns of functional connectivity. The extent to which this apparent transition in functional connectivity relates to other ongoing neurodevelopmental changes is an open question. Development of white matter tracts is a likely contributor to changes in functional connectivity. White matter volume demonstrates ongoing increase over middle to late childhood (Giedd et al. 1999); this is at least partly accounted for by myelination of axons, which continues throughout this period (Yakovlev and Lecours 1967). Developmental increases in white matter integrity (as measured by fractional anisotropy) vary as a function of tract; of particular note, the superior longitudinal fasciculus which serves as the major conduit of frontoparietal communication, demonstrates ongoing increase in integrity through age 18. Thus, ongoing changes in white matter might enable the observed changes in functional connectivity.

In addition to changes in long-range fiber tracts, it is possible that synaptic changes could also affect developing patterns of functional connectivity. Large-scale synaptogenesis—the creation of new synaptic connections—is largely complete by early childhood (Rakic et al. 1994; Petanjek et al. 2008). Synaptic density plateaus during childhood and then begins a sharp decline with the onset of synaptic pruning. Notably, there are large regional differences in the timing of this transition, and PFC stands out for the late (mid-adolescent) onset and extended duration of large-scale pruning (Huttenlocher and Dabholkar 1997; Petanjek et al. 2011; Liu et al. 2012). Thus, overall changes in synaptic density are limited during middle to late childhood. However, synaptic reorganization and turnover continues throughout this period (Rakic et al. 1994; Selemon 2013), and could play a role in the changes in functional connectivity observed here.

Finally, there is evidence that hormonal changes that occur during puberty can affect both structural and functional connectivity (Peper et al. 2011). Hormonal changes, in particular, may help to explain the timing of the changes that we observed here.

Despite our efforts to mitigate the effects of motion on our results, findings related to age-related differences in functional connectivity must be interpreted with caution. The observed increases in fronto-parietal connectivity and decreases in intrafrontal and intraparietal connectivity fit into the pattern—well established in the literature (Fair et al. 2009; Supekar et al. 2009) —of age-related increases in long-range connectivity and decreases in short-range connectivity. But these also conform to the changes that one would expect due to decreases in head motion (Power et al. 2012; Satterthwaite et al. 2012; Van Dijk et al. 2012). Arguing against the idea that head motion could account for these results is the fact that we observe different patterns of developmental change among the shorter range connections that we examined and also among the longer range connections. These results highlight the importance of examining specific connections, instead of or in addition to looking at general classes of connections such as long range versus short range.

Also relevant to the interpretation of age-related changes is the fact that longitudinal attrition differed as a function of SES. Lower SES participants were less likely to return for a second scanning session, and so it is in theory possible that mixedmodel results are biased toward changes that are seen in higher SES individuals. However, all of the effects that we obtained via mixed-model analysis with longitudinal data were also present when the analyses were restricted to cross-sectional data.

Fronto-parietal Connectivity and Reasoning Ability

In adults, RLPFC and IPL are 2 regions that are consistently engaged by higher order relational reasoning tasks. Thus, our main hypothesis concerned functional connectivity between RLPFC and IPL. Specifically, we hypothesized that connectivity between these regions should be an important predictor of reasoning ability. We considered that this might be the case regardless of maturity, or that it might only be true as the reasoning system approaches its adult state. In fact, we observed the latter pattern: left RLPFC-IPL connectivity was the strongest predictor of reasoning ability, but only in older children and adolescents. This result confirms the importance of RLPFC-IPL connectivity in the mature reasoning system, but also indicates that there are meaningful differences in the way that reasoning is performed in an immature neural system. It is an open question whether these differences reflect operation of different brain mechanisms in support of reasoning in younger versus older children, or if they reflect differential importance of the same set of mechanisms.

Prior investigations of the link between developing functional connectivity and higher cognition have also pointed to the importance of the fronto-parietal connection. For example, Ezekiel et al. (2013) demonstrated increased connectivity of lateral PFC to the anterior cingulate cortex (ACC) and to IPL during execution of the dimensional change card sorting task, in adults relative to children. Langeslag et al. (2012) demonstrated a relationship between nonverbal intelligence and increased functional connectivity between right parietal and frontal regions and between right parietal and dorsal ACC. Emerson and Cantlon (2012) reported a positive relationship between fronto-parietal connectivity and children's math ability. Complementing these developmental findings, Mackey et al. (2013) showed that intensive reasoning training in young adults is associated with increased RLPFC-parietal connectivity, with no change in DLPFC-parietal connectivity. Thus, our results add to the general finding that fronto-parietal connections can be critical for many higher cognitive tasks.

Notably, we did not observe a relation between average network connectivity and reasoning ability. Moreover, we did not observe a consistent relation between average fronto-parietal, or even RLPFC-parietal, connectivity and reasoning ability. Instead, we observed a very specific pattern: reasoning ability was positively related to RLPFC-IPL connectivity but negatively related to RLPFC-SPL connectivity. These results contrast with the results from prior studies of functional connectivity and general intelligence, which have linked IQ to generalized increases in functional connectivity (Li et al. 2009; van den Heuvel et al. 2009; Cole et al. 2012; Langer et al. 2012). The present results suggest that more specific links between IQ and functional connectivity might be found. On the other hand, just as reasoning ability is a specific component of IQ, connectivity changes associated with improvements in reasoning may be more specific than changes associated with more general intellectual improvement.

An Emerging Network for Reasoning

Beyond the hypothesized relationship between RLPFC-IPL connectivity and reasoning that was present in older children, we observed a compelling pattern of relations between functional connectivity and reasoning ability in the younger and middle child groups. We have demonstrated previously the occurrence of a developmental shift, from early childhood to adolescence, in the brain regions that are most engaged by higher order relational reasoning tasks (Wendelken et al. 2011). Specifically, while selective engagement for higher order reasoning is limited to DLPFC in younger children, it shifts to include RLPFC and IPL in adolescents and adults. Here, we observed a similar developmental shift in functional connectivity. In the youngest group of children, the factor that most related to reasoning ability was the connectivity of DLPFC and VLPFC. This was a negative relationship, with decreased connectivity associated (albeit weakly) with improved reasoning ability. Thus, at a relatively early stage in the development of the reasoning system, a key neurodevelopmental change may be the differentiation of DLPFC-the region that is most engaged by reasoning tasks in this age range-from nearby VLPFC. In the middle group of children, further differentiation of DLPFC from VLPFC appears less important for reasoning, perhaps because it is mainly complete or because of the shift toward a more prominent role for RLPFC. Instead, in this group, it was tighter coupling of left and right RLPFC that was most closely associated with reasoning ability. This pattern, along with the emerging importance of the RLPFC-IPL connection, reinforces the idea of an anterior shift within lateral PFC for higher order reasoning.

Processing Speed and Working Memory

Behavioral studies measuring concurrent and longitudinal relationships between cognitive abilities have pointed to processing speed and working memory as key mediators of reasoning development (e.g., Kail and Salthouse 1994; Li et al. 2004; Nettlebeck and Burns 2010; Demetriou et al. 2014). Large improvements in reasoning ability during the early school years, observed here as well as in prior studies (Goswami 1991; Ferrer and McArdle 2004), are complemented by large improvements in both processing speed and working memory (Fry and Hale 1996). In recent work, we demonstrated that processing speed mediates the concurrent relationship between global white matter integrity and reasoning across childhood (Ferrer et al. 2013). Thus, we reasoned that processing speed and/or working memory might similarly mediate the concurrent relationship between functional connectivity and reasoning ability.

As in prior behavioral work, processing speed was related to reasoning ability in our sample, particularly in the youngest group of children. However, processing speed did not mediate any of the observed effects of functional connectivity on reasoning ability. This finding reveals a difference between structural and functional connectivity. On one hand, as we have shown previously, the widespread increase in white matter coherence observed during development contributes to reasoning ability indirectly via its influence on processing speed. On the other hand, developmental changes (both increases and decreases) in functional connectivity between specific nodes in the lateral fronto-parietal network contribute to reasoning ability in a manner that cannot be explained by improved processing speed. In older children and adolescents, the relationship between fronto-parietal connectivity and reasoning ability was not mediated by either processing speed or working memory. In middle children, in contrast, working memory did mediate the relationship between intraprefrontal connectivity and reasoning ability. In younger children, it was neither functional connectivity nor working memory, but rather processing speed, that had the strongest relation to reasoning ability. Thus, our results are consistent with a developmental cascade model of the relationship between processing speed, working memory, and reasoning ability (Fry and Hale 1996). Specifically, they indicate that development of reasoning ability is driven initially by improvements in processing speed and subsequently by improvements in working memory.

Summary

In previous work, we have argued that neurodevelopment of the reasoning system is characterized by shifting patterns of activation, particularly within lateral PFC. Here, we provide evidence that patterns of functional connectivity undergo a similar neurodevelopmental change. In particular, there is a shift in the importance of functional connections for reasoning toward connections that involve RLPFC, and ultimately toward specificity of RLPFC-parietal communication. This shift occurs alongside developmental changes in the relationships between reasoning, processing speed, and working memory. Much work remains to be done to understand how the specific brain regions and their interactions contribute to reasoning and its development. However, knowing which connections matter, and when, is a critical step toward a deeper mechanistic understanding.

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