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Morphometry of left frontal and temporal poles predicts analogical reasoning abilities

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Keywords: analogy, frontal pole, morphometry, reasoning, rostral prefrontal cortex

Abstract

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10 Analogical reasoning is critical for making inferences and adapting to novelty. It can be studied
11 experimentally using tasks that require creating similarities between situations or concepts, i.e.,
12 when their constituent elements share a similar organization or structure. Brain correlates of
13 analogical reasoning have mostly been explored using functional imaging that has highlighted
14 the involvement of the left rostrolateral prefrontal cortex (rLPFC) in healthy subjects. However,
15 whether inter-individual variability in analogical reasoning ability in a healthy adult population is
16 related to differences in brain architecture is unknown. We investigated this question by
17 employing linear regression models of performance in analogy tasks and voxel-based
18 morphometry in 54 healthy subjects. Our results revealed that the ability to reason by analogy
19 was associated with structural variability in the left rLPFC and the anterior part of the
20 inferolateral temporal cortex. Tractography of diffusion-weighted images suggested that these
21 two regions have a different set of connections but may exchange information via the arcuate
22 fasciculus. These results suggest that enhanced integrative and semantic abilities supported by
23 structural variation in these areas (or their connectivity) may lead to more efficient analogical
24 reasoning.
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3 *“When a man sits with a pretty girl for an hour, it seems like a minute. But let him sit on a hot*
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6 *stove for a minute and it's longer than any hour. That's relativity”* (Einstein 1938).
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9 This metaphorical quote from Albert Einstein illustrates how an analogy can be
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11 employed to make the concept of relativity a little more comprehensible to the layman. To
12
13 understand this metaphor, we employ analogical reasoning to identify similarities between a
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15 familiar situation (variations in time estimation according to the pleasantness of a personal
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17 experience) and a new or complex one (variation in space and time according to the referential of
18
19 the observant). Analogical reasoning uses these similarities to make inferences about a novel
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21 situation or concept and to infer rules and implications (Gentner and Holyoak 1997). Analogical
22
23 reasoning is therefore critical for adapting to novelty, and for learning, explaining or conceiving
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25 new concepts, and is thought to constitute a cognitive basis for fluid intelligence (Holyoak and
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27 Thagard 1995; Geake and Hansen 2005; Hofstadter and Sander 2013).
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32 Analogical reasoning is considered a form of relational reasoning, as similarities concern
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34 the relationships between the elements of a situation or an object rather than these elements
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36 themselves (Blanchette and Dunbar 2000; Christoff et al. 2009; Markman and Gentner 2000).
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38 These relationships describe particular aspects of the “structure” of an object/situation (or how
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40 the elements are organized in an object/situation). Cognitive theories assume that analogy
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42 processing includes the generation of mental representations of the relationships (their structure)
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44 and their mapping based on their similarities, which is distinct from the mapping of stimuli based
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46 on non-relational item-to-item similarity (Blanchette and Dunbar 2000; Holyoak and Morrison
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48 2012; Holyoak and Thagard 1995; Markman and Gentner 2000; Markman and Gentner 1993).
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50 Processes allow for the generation of a schema for the whole analogy, i.e., a general relational
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3 concept of the common structure. In this sense, analogical reasoning includes a conceptual
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5 dimension that is not present in purely perceptual similarity matching.
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9 An analogy schema can be inferred from the mapping of several exemplars sharing a
10 similar structure (exemplar-based). For instance, in two sets of three different letters, an analogy
11 schema (i.e., a relational concept) can be produced through a similarity between sets that show a
12 similar pattern of 'increase in size', i.e., by noting that in both sets the first letter is smaller than
13 the second one and that the second one is smaller than the third one. Alternatively, when the
14 schema is familiar, it can be retrieved from memory based on only one familiar exemplar or on a
15 verbal description (e.g. 'increase in size'), i.e. a relational term describing the conceptual analogy
16 (concept-based) (Gentner and Medina 1998). An exemplar-based analogy requires the induction
17 of the analogy schema, whereas in concept-based analogy the analogy schema is given with the
18 instruction. Whether exemplar-based and concept-based analogical reasoning processing is
19 supported by different brain structures is unknown. Studies of inductive reasoning nevertheless
20 suggest that the left lateral prefrontal cortex is important for inferring abstract rules (Reverberi et
21 al. 2005a; 2005b), such as analogy schemas.
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40 Functional imaging studies of analogical reasoning have shown a set of frontal and
41 parietal regions engaged during analogy tasks (Wharton et al. 2000; Christoff et al. 2003; Bunge
42 et al. 2005, 2009; Geake and Hansen 2005, 2010; Green et al. 2006, 2010, 2012; Cho et al. 2010;
43 Wendelken et al. 2008, 2012; Krawczyk et al. 2010a; Volle et al. 2010; for a review Krawczyk
44 2012). The fronto-parietal network involved in analogical reasoning has been associated with the
45 executive or working memory aspects of analogy tasks and with fluid reasoning (Jung and Haier
46 2007; Geake and Hansen 2010; Preusse et al. 2011). The parietal component may be involved in
47 processing the visuospatial relationships between multiple objects (Watson et al. 2012) or in the
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3 organization of maintained information (Wendelken et al. 2008). Caudal prefrontal regions may
4 support the inhibition of interference (Morrison et al. 2004, Krawczyk et al. 2008; Cho et al.
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6 2010; Thibaut et al. 2010b), or the controlled retrieval or selection of information in semantic
7
8 memory (Bunge et al., 2005). A few patient studies have demonstrated the critical role of the
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10 PFC in analogical reasoning (Krawczyk et al., 2008; Morrison et al., 2004) and additionally
11
12 suggested that the temporal cortex may play a significant role in analogical reasoning by
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14 activating the semantic relation that links the terms of the analogy (Morrison et al., 2004;
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16 Schmidt et al., 2012). Among the regions that have been associated with analogical reasoning,
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18 the left rostrolateral prefrontal cortex (rlPFC) is the most consistently activated in functional
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20 imaging, as demonstrated in a recent meta-analysis of 10 functional imaging studies on analogy
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22 (Vartanian 2012). The rostral PFC component is thought to support the simultaneous comparison
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24 and/or integration of multiple relations between stimuli, the integration of the results of separate
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26 cognitive operations (Ramnani and Owen 2004; Christoff et al. 2001; Kroger et al. 2002;
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28 Wendelken et al. 2008; Bunge et al. 2005; 2009; Cho et al. 2010; Hampshire et al. 2011), or high
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30 levels of abstract thinking (Christoff et al. 2009). Previous works have also suggested that the
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32 rostral PFC may be recruited before exemplars are compared (Volle et al. 2010; Krawczyk et al.
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34 2010a), bringing its role in exemplar-based analogy into question. The exact role of this region in
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36 distinct analogy processes such as integration of relationships, mapping, schema induction, and
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38 its relationship to analogy performance remain unclear.
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49 Analogical reasoning is indeed a complex and highly adaptive cognitive function marked
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51 with a large variability in individual ability. Studies in children and adolescents previously
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53 suggested that the development of analogical reasoning abilities with age is related to executive
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55 functions (Thibaut et al. 2010a; 2010b) and to changes in rlPFC activation and structure (Crone
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3 et al. 2009; Dumontheil et al. 2010; Krawczyk et al. 2010b). Although evidence from activation
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5 studies is increasing in adults, the relationship between the morphology of the PFC (or other
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7 brain regions) and an individual's capacity to reason by analogy remains unexplored. In other
8
9 words, can inter-individual variations in analogy ability be linked to individual variations in
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11 regional structures of the brain?
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16 Brain local morphology is classically measured using voxel-based morphometry (VBM;
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18 Ashburner and Friston 2000; Good et al. 2001; Kanai and Rees 2011) and can be statistically
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20 related to behavioral measures. Previous morphometry studies have shown correlations between
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22 individual performance in other relational reasoning tests, such as IQ tests reflecting fluid
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24 intelligence, and grey matter (GM) local morphology in the rLPFC (Frangou et al. 2004; Haier et
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26 al. 2004; Colom et al. 2006, 2009; Narr et al. 2007; Yuan et al. 2012), and between analogical
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28 reasoning capacity and the structure of the developing brain, but have not explored analogical
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30 reasoning in adults. Whether rLPFC structural variability in adults relates to variable performance
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32 in analogical reasoning (a particular type of relational reasoning) and to the processes engaged
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34 when forming the analogy remains to be demonstrated.
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41 To address this question, we performed a voxel-wise analysis to correlate local brain
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43 volumes with analogical reasoning performance. To distinguish between the relational
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45 processing component and the schema induction component of analogical reasoning, we
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47 manipulated our analogy tasks to distinguish analogies based on exemplar comparison
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49 (exemplar-based analogies requiring schema induction) from analogical reasoning based on a
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51 relational term (concept-based analogies in which an analogy schema is provided to the
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53 participants, for instance 'increase', without a need to infer it). We used a control-matching task
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55 based on the similarity of perceptual features in order to control for non-relational mapping
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3 processing. We explored the anatomical connectivity of the regions associated with these tasks
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5 using diffusion-based tractography to provide further understanding about their potential
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7 interactions and roles in analogy processing.
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10 11 12 13 **Methods**

14 15 *Participants*

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17 Fifty-seven volunteers were initially included in this study, but three of them were excluded from
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19 the analysis for medical reasons (anomalies on neuropsychological testing or on the brain MRI).
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21 Thus, fifty-four right-handed native French speakers (27 females; age 22 - 71 years, mean $45.8 \pm$
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23 14.4 years) participated in the morphometry study. A large age range was chosen in order to
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25 include a group of unselected participants with enough variability to represent the human
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27 diversity in the general population. The advantages of such an approach have been discussed
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29 previously (Colom et al., 2007; Haier et al., 2004; Goh et al., 2011; Grogan et al., 2009).
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31 Neuropsychological and radiological data were carefully screened. All participants were healthy
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33 adults with no history of neurological or psychiatric disorders and no cognitive impairments or
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35 depression, as assessed using translated versions of the Mini Mental state (Folstein et al. 1975)
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37 and the MADRS (Montgomery-Asberg Depression Scale; Montgomery and Asberg 1979) as
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39 well as the Frontal Assessment battery (Dubois et al. 2000). All brain images were examined by
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41 a neuroradiologist. Millimetric T1-weighted and diffusion weighted images showed no
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43 significant signal abnormalities evocative of a small vessel disease or of an evolving
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45 neurological disease. Subjects with MRI pathological abnormalities were excluded. Participants
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47 had an average of 15.4 ± 3.0 years of education (range 10 - 26).
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3 The local ethics committee approved the experiment, and all participants provided written
4 informed consent.
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8 Of the 54 participants, 47 (24 males; mean age 45.5 ± 14.8 years; mean education level
9 15.4 ± 3 years) were included in the connectivity study using diffusion images. Data from the
10 remaining 7 participants could not be analyzed because of technical problems with the diffusion
11 images.
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17 18 19 20 21 ***Experimental Procedure (Figure 1)*** 22

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24 The experimental paradigm consisted of four experimental conditions: two tasks
25 (Analogy and Match) each having an exemplar-based condition ('Find') and a concept-based or
26 rule-based condition ('Apply') as described below. The participants were trained on the two
27 analogy (named AnalogyApply and AnalogyFind) and two match (MatchApply and MatchFind)
28 conditions for 26 trials. All subjects understood the instructions and were able to perform the
29 tasks correctly after the training. Then, each condition was implemented in blocks in the
30 following order: 28 MatchApply, 28 MatchFind, 48 AnalogyApply and 48 AnalogyFind trials.
31 The trial order was randomized within each block. In each trial, instructions were displayed for
32 four seconds. Immediately afterward, a left set of stimuli (source set) was displayed for 2
33 seconds followed by the display of the two comparison sets on the right (target sets). The
34 participants were required to judge which of the two target sets shared similarities with the
35 source set on criteria indicated by the instructions of the task condition. The source and target
36 sets consisted of groups of letters, numbers, or abstract symbols in different colors, sizes, or
37 patterns. The sets were equivalent in terms of visual and temporal features between the task
38 conditions. The participants had 11.5 seconds to respond by pressing the up or down arrow key.
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3 A signal appeared 1.5 seconds before the end of the display. Feedback was displayed for 0.5
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5 seconds, a green circle indicated a correct answer and a red circle an incorrect answer. The trials
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7 were separated by a 5-second interval.
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10 11 12 13 14 ***Behavioral Task (Figure 1)*** 15

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17 The Analogy and control Match tasks employed were adapted from a previous study
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19 (Volle et al. 2010). The participants had to choose the target sets that shared similarities with the
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21 source set, based on the relationships between the stimuli (the schema; Analogy task), or based
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23 on a shared perceptual attribute (Match task). Hence, this experimental design allowed for the
24
25 study of the participants' performance of finding or applying an abstract analogy schema
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27 (concept) while controlling for their performance in finding or applying a perceptual (concrete)
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29 similarity.
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34 Six diverse attributes were used as matching rules in the Match task, namely color,
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36 quantity, size, texture, figures and letters. In the Analogy task, 6 distinct schemata were
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38 employed, namely proportion, subtraction, addition, mirroring, symmetry and progression (see
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40 supplementary Figure S1 for a sample of trials). As described in Volle et al. (2010) and
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42 illustrated in supplementary Figure S1, half of the Analogy trials were intra-dimension analogies,
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44 and half were cross-dimension analogies. In the intra-dimension task, the analogy schema
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46 concerned the same dimension in the source and target sets (e.g., an increase in the size of the
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48 stimuli in both the source and the target). In the cross-dimension task, the analogy concerned
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50 different dimensions (for instance, an increase in the size of the stimuli in the source and an
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52 increase in the color lightness of the target stimuli).
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Original Analogy and Match tasks were further modified in order to differentiate exemplar-based and concept-based analogical processing. In the original exemplar-based Analogy condition (as used in Volle et al. 2010, here named AnalogyFind), participants had to find the relational schema by considering the structures of each exemplar set, comparing them and finding their similarities (i.e., it was an internally generated analogy with relational processing and concept formation/induction). The instruction “find analogy” was displayed, and the task was solved by comparing the sets (Figure 1, top left). In the new concept-based Analogy task, here named AnalogyApply condition, the relational schema was explicitly given to the subjects using a verbal term displayed on the screen (as for instance “mirror image” in Figure 1, bottom left). They had to consider and compare the multiple relationships between stimuli, but there was no need to form or retrieve the schema (i.e., it was an externally driven analogy with relational processing but without schema induction). The three analogy schemas used in the AnalogyFind condition, were distinct from the three schemas used in the AnalogyApply condition.

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The same principle was applied to the Match tasks. In the MatchFind condition, participants had to find the perceptual relationship between the source and correct target set. The instruction “find match” was displayed and the task was solved by comparing the exemplars and finding the matching rule (Figure 1, top right, the similarity concerned the number of stimuli). In the MatchApply condition, the participants were instructed to apply a given matching rule, which was presented verbally. For instance, in Figure 1 (bottom right), “Same Colors” appeared on the screen to instruct the participants to match colors. The three matching rules used in the MatchFind condition, were distinct from the three rules used in the MatchApply condition.

Behavioral Analysis

Accuracy and response times were measured, and statistical analyses were conducted. Repeated measures ANOVA analysis was employed to compare conditions in the 2x2 within-subjects design for the Analogy versus Match task and Find versus Apply conditions using SPSS software (<http://www-01.ibm.com/software/analytics/spss/>). We also ran Pearson correlation analyses between the age, education, and experimental scores, and compared the performance of males and females using an independent samples t test.

VBM study: Image acquisition and analysis

Structural T1-weighted images

All participants underwent the same high-resolution T1-weighted structural MRI scans acquired on a Siemens 3 Tesla VERIO TIM system equipped with a 32-channel head coil. An axial three-dimensional MPRAGE dataset covering the whole head was acquired for each participant as follows: 176 slices, voxel resolution = $1 \times 1 \times 1$ mm, TE = 2.98 msec, TR = 2300 msec, flip angle = 9° .

VBM pre-processing

3D T1-weighted sequences were processed and analyzed with SPM8 (Wellcome Department of Imaging Neuroscience, London, UK) running on Matlab (Mathworks Inc., Natick, MatchApply, USA; www.mathworks.com/matlabcentral). The VBM8 toolbox (<http://dbm.neuro.uni-jena.de/vbm/>) was employed to perform MRI data pre-processing (<http://dbm.neuro.uni-jena.de/vbm8/BVM8-Manual.pdf>). First, the T1 images were spatially normalized to the

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2
3 MNI152 Dartel template using the high-dimensional Dartel normalization (Ashburner 2007) and
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5 were segmented into GM, WM and cerebrospinal fluid using SPM8's new version of the unified
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7 segmentation method (new segment; Ashburner and Friston 2005). Default estimation
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9 parameters were employed (<http://dbm.neuro.uni-jena.de/vbm8/BVM8-Manual.pdf>) to compute
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11 normalized and modulated GM images with an isotropic voxel size of 1.5 mm³. Modulation
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13 compensates for regional volume changes caused by normalization. The “normalized non-linear
14
15 modulation only” option was used, allowing us to analyze relative differences in regional GM
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17 volume corrected for individual brain size. The quality was evaluated by displaying one slice for
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19 each image module and searching for visual abnormalities and by checking the sample
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21 homogeneity using the covariance between individual images. The images with the lowest
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23 covariance (-2 standard deviations) were visually examined, but none of them had to be
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25 excluded. In addition, all normalized 3D images were visually inspected and compared to the
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27 template using frontal anatomical landmarks by an expert neurologist (E.V.). Modulated and
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29 normalized GM images were then smoothed using a Gaussian kernel of 8-mm³ full width half
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31 maximum (FWHM) to account for slight variations between individual normalizations and to
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33 allow for parametric statistics. After pre-processing, the smoothed, modulated, normalized GM
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35 datasets were used for statistical analyses.
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46 VBM whole-brain statistical analysis

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49 To investigate the relationship between VBM regional GM density and various aspects of
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51 analogical reasoning, we ran multiple regression analyses in SPM8 between GM volume and
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53 Analogy and Match mean scores. Two separate models were used. In the first one, the Find and
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55 Apply conditions were averaged so that the mean Analogy score (AnalogyFind and
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3 AnalogyApply) and the mean Match score (MatchFind and MatchApply) were entered as
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5 separate covariates in the regression model, enabling the determination of the brain correlates for
6
7 each task. In the second model, the Analogy and Match scores were collapsed so that the mean
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9 score in AnalogyFind and MatchFind conditions (“Find”) and the mean score in AnalogyApply
10
11 and MatchApply conditions (“Apply”) were entered as separate covariates in the regression
12
13 model. Age, gender and education were co-varied out in the linear regression model. Data were
14
15 also normalized and corrected for individual total GM volume by entering their values as
16
17 covariates in the linear model. Global values of total GM volume were extracted and calculated
18
19 from the `get_totals` script (available on
20
21 http://www0.cs.ucl.ac.uk/staff/g.ridgway/vbm/get_totals.m). Threshold masking was set to 0.2 to
22
23 include in the analysis only voxels with sufficient signal. For each regression analysis, we
24
25 investigated significant results at $p < 0.001$ uncorrected for multiple comparisons (with a
26
27 minimal cluster size of 100 voxels). Within the significant clusters, the mean GM volume was
28
29 extracted using FSL (<http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/>) and entered as dependent variables in
30
31 new regression analyses, in which each task condition was a covariate, and age, gender,
32
33 education and total GM volume were covaried out. Each cluster volume was plotted against
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35 performance for illustration purposes.
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44 Next, a *Small Volume Correction* (SVC) for multiple comparisons was applied on the
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46 same analyses as described in the following paragraph. This statistical approach of combining
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48 uncorrected and corrected results is frequently employed in VBM analysis (Ridgway et al. 2008)
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50 and has the advantage of showing both exploratory and hypothesis driven results.
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57 VBM SVC Analyses within Independent Regions of Interest (ROI)

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3 ROIs were selected independently from the whole-brain analysis and located in the most
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5 consistent region reported in functional imaging studies: the rIPFC. The left ROI was based on
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7 the previous functional brain imaging study by Volle and colleagues (Volle et al. 2010) that
8
9 demonstrated rostral prefrontal involvement in analogical reasoning when employing a similar
10
11 task. The analysis was focused on the rIPFC maxima reported in this study of analogy tasks. The
12
13 ROI was located in the left rostral MFG or BA 10/46 (MNI coordinates: $x=-44$, $y=50$, $z=-4$). We
14
15 built an 8-mm radius sphere centered on these coordinates and used this ROI for subsequent
16
17 analyses. Note that this region is very similar to the region reported in the meta-analysis by
18
19 Vartanian (Vartanian, 2012; MNI-converted coordinates, $x=-44.4$, $y=40.1$, $z=3.2$) and identified
20
21 as a core region for analogy. To check if analogy processing is associated with bilateral rIPFC
22
23 morphometry, we built a symmetrical ROI in the right hemisphere with the following
24
25 coordinates ($+44$, 50 , -4). We ran SVC analyses in these ROIs to investigate significant
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27 correlations of each condition with GM volume within these regions. The threshold was set to
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29 0.05 after a *Family Wise Error* correction (fwe) for multiple comparisons.
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40 ***Connectivity study: image acquisition, preprocessing and analysis***

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42 One of the best methods of studying the functional specialization of specific brain regions is to
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44 examine the input and output of that region (Van Essen and Maunsell 1983). Therefore, based on
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46 the diffusion obtained from 47 of the 54 participants, we explored the connections terminating in
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48 and emerging from the brain regions identified by VBM as showing a volumetric change
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50 associated with performance in analogical reasoning.
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Diffusion images acquisition

A total of 70 near-axial slices were acquired on a Siemens 3-Tesla VERIO TIM system equipped with a 32-channel head coil. We used an acquisition sequence fully optimized for tractography of DWI that provided isotropic ($2 \times 2 \times 2$ mm) resolution and coverage of the whole head. The acquisition was peripherally gated to the cardiac cycle with an echo time (TE) of 85 msec. We used a repetition time (TR) equivalent to 24 RR. At each slice location, 6 images were acquired with no diffusion gradient applied. Additionally, 60 diffusion-weighted images were acquired in which gradient directions were uniformly distributed in space. The diffusion weighting was equal to a b-value of 1500 sec/mm^2 .

Diffusion imaging pre-processing

Spherical deconvolution was chosen to estimate multiple orientations in voxels containing different populations of crossing fibers (Tournier et al. 2004; Anderson 2005; Alexander 2006). The damped version of the Richardson-Lucy algorithm for spherical deconvolution (Dell'Acqua et al. 2010) was calculated using an in-house developed software. Algorithm parameters were chosen as described before (Dell'Acqua et al. 2013).

Whole-brain tractography was performed by selecting every brain voxel with at least one fiber orientation as a seed voxel. From these voxels and for each fiber, orientation streamlines were propagated using Euler integration with a step size of 1 mm. When entering a region with crossing white matter bundles, the algorithm followed the orientation vector of the least curvature (as described in Schmahmann et al. 2007). Streamlines were halted when a voxel without fiber orientation was reached or when the curvature between two steps exceeded a

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3 threshold of 60°. Spherical deconvolution, fiber orientation vector estimation and tractography
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5 were performed using in-house software developed with Matlab 7.8
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8 (<http://www.mathworks.com>).
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11 12 13 Tractography dissections

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16 Significant regions from the whole-brain VBM analyses were used as ROIs for tract dissections.
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18 We dissected the tracts connecting the observed ROIs associated with Analogy, Match and Find
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20 performance.
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24 For each participant, the convergence speed maps (Dell'Acqua et al. 2013) were
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26 registered to the MNI152 template using Advanced Normalization Tools ANTs (Klein et al.
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28 2009). The inverse deformation was then applied to the ROIs to bring them within the native
29
30 space of every participant.
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34 Binary individual visitation maps were created for the connections emerging from or
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36 terminating in the three observed ROIs by assigning each voxel a value of 1 or 0, depending on
37
38 whether the voxel was intersected by the streamlines of the tract. Binary visitation maps of each
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40 of the dissected tracts were normalized to MNI space using the same affine and diffeomorphic
41
42 deformations as calculated above. We created percentage overlap maps by adding the normalized
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44 visitation maps from each subject at each point in the MNI space. Therefore, the overlap of the
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46 visitation maps varies according to inter-subject variability. We inspected tracts reproducible in
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48 more than 50% of the participants, a method described previously in Thiebaut de Schotten et al.
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50 (2011). Tracts resulting from this analysis were visually inspected and identified using an atlas of
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52 human brain connections (Rojkova et al. under revision; Thiebaut de Schotten et al. 2011).
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Results

Behavioral Results

Accuracy (Figure 2A)

Descriptive statistics revealed systematic errors in a few trials. Less than 50% of the participants (less than chance) gave correct answers to three of the 48 AnalogyApply trials and three of the 48 AnalogyFind trials. Because these trials could have been missed for other reasons than a poor analogical reasoning ability, or because in these trials both target sets could be interpreted as a correct analogy, they were discarded from further statistical analyses, and only the remaining 45 AnalogyApply and 45 AnalogyFind trials were analyzed. Repeated measures ANOVA revealed a significant main effect of task ($F(1, 53) = 70.1, p < .001$; Match conditions mean = 93.7% of correct responses; Analogy conditions mean = 85.4%) and a marginally significant main effect of concept formation (finding the schema; $F(1, 53) = 3.7, p = .06$; Apply conditions mean = 89.5%; Find conditions mean = 87.7%). No significant interaction was found between task and concept formation effects, $F(1, 53) = 0.7, p < .401$. There was no decrease in performance over time in our group of participants.

Response Times (RTs; Figure 2B)

Repeated measures ANOVA revealed a significant main effect of task on RTs ($F(1, 53) = 157.4, p < .001$; Match conditions mean = 3347 ms; Analogy conditions mean = 4397 ms) and a significant main effect of concept formation ($F(1, 53) = 87.0, p < .001$; Apply conditions mean

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3 in Analogy tasks (mean score of AnalogyApply and AnalogyFind conditions) was associated
4 with GM volume in the left and right anterior inferolateral temporal cortex (aITG; regions
5 hereafter referred to as lTEMP on the left and rTEMP on the right), while accuracy in Match
6 tasks (mean score of MatchApply and MatchFind conditions) was not associated with a
7 significant region (Figure 3).
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16 Negative GM correlations are reported in Table 2. For accuracy in Analogy tasks,
17 negative correlations with GM volume were found in the left rIPFC (middle frontal gyrus
18 (MFG); BA 10; a region hereafter referred to as lPOL). For accuracy in Match tasks, negative
19 correlations with GM volume were observed within the anterior part of the right inferomedial
20 temporal cortex (fusiform gyrus) (region hereafter referred to as rTEMPmatch) and in the
21 parietal region.
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30 Because age was negatively correlated with performance in Analogy tasks, and because
31 the age range in our participants was large, we searched for correlations between age and GM
32 volume within clusters associated with Analogy performance in the whole-brain analysis. The
33 correlation between age and GM volume within the left rIPFC region (lPOL) was significant and
34 negative ($r = -.435$; $p = 0.001$), but it was no longer significant after controlling for the total GM
35 volume ($r = -.202$, $p = .147$). Within the anterior temporal region (lTEMP, that correlated
36 positively with analogy accuracy), the correlation between age and GM volume was significant
37 and negative ($r = -.296$, $p = .030$), but was not significant when controlling for total GM volume
38 ($r = -.161$, $p = .250$).
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55 ***VBM Whole-Brain Analysis: GM Correlations with “finding the rule” (Figure 3)***
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Negative GM correlations are reported in Table 2. For accuracy in the Find conditions (mean AnalogyFind and MatchFind accuracy), negative correlations with GM volume were found in the left inferior frontal sulcus (IFS; BA 45) extending to the pars triangularis and the middle frontal gyrus; a region hereafter referred to as lmidPFC. No significant correlation was found with the Apply conditions.

The correlation between age and GM volume within the lmidPFC region was significant and negative ($r = -.642$; $p < .001$), and stayed significant after controlling for the total GM volume ($r = -.388$, $p = .004$).

Regression with the VBM regions (Table 3 and Figure 3)

To better understand the brain correlates to each condition and task, we ran multiple regressions between each region identified in the whole-brain analysis (as dependent variables) and each score separately (AnalogyFind, AnalogyApply, MatchFind and MatchApply being predictors), with age, gender, education and total GM volume as covariates. The plots are presented in Figure 3 and the statistics in Table 3. These analyses show that the Analogy regions (lPOL, rTEMP) are associated with both the AnalogyFind and AnalogyApply scores, and that the Find region (lmidPFC) is associated with both the AnalogyFind and MatchFind scores.

VBM SVC Analysis (Table 4)

To clarify the role of rIPFC in analogy, we looked for significant correlations between GM volume in rIPFC ROIs with mean Analogy and mean Find performance (as in the whole-brain analysis), and additionally with each experimental condition in separate regression models. Left

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3 and right rLPFC ROIs have been drawn *a priori* from fMRI data (Volle et al. 2010). Significant
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5 negative correlations were found for accuracy in the AnalogyFind and AnalogyApply conditions
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7 and for the mean Analogy score (Table 4) within the left rLPFC ROI, but not in the right rLPFC.
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9 For MatchApply and MatchFind conditions and for the Find mean score, no correlation was
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11 found within any ROI.
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18 **Connectivity patterns of the VBM regions (Figure 4)**

19 ***Connectome of Analogy regions***

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22 The IPOL connectome, representing fibers connecting the left rLPFC VBM region associated
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24 with analogy performance, included projection fibers from the anterior fronto-thalamic
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26 radiations, commissural fibers from the anterior forceps of the corpus callosum, and several
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28 association fibers, namely the Inferior Fronto-Occipital fasciculus (IFOF), the Uncinate
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30 fasciculus (UF), the Arcuate fasciculus (AF, long segment) and the Fronto-Marginal tract
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32 (FMT).
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39 The ITEMP connectome, representing fibers connecting the left temporal VBM region,
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41 was identified as the Inferior longitudinal fasciculus (ILF), the Arcuate fasciculus (AF, long
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43 segment), and the tapetum of the corpus callosum. The rTEMP connectome included similar
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45 contralateral fasciculi in the right hemisphere.
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49 These findings show that IPOL and ITEMP have distinct anatomical connections but are
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51 both connected to the long segment of the AF.
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57 ***Connectome of Match (control task) region (Figure 4)***

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3 The rTEMPmatch connectome (representing fibers connecting the right inferomedial temporal
4 VBM region associated with Match tasks) included the ILF (running medially to the rTEMP
5 connectome), the fornix, and intratemporal connections.
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10 11 12 13 14 ***Connectome of Find regions (Figure 4)*** 15

16 The lmidPFC connectome (representing fibers connecting the left lateral caudal prefrontal VBM
17 region associated with finding the matching rule and/or the analogy schema) included mainly
18 intrafrontal fibers along the IFS and inferior frontal gyrus, and possibly some arcuate fibers
19 posteriorly.
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30 **Discussion** 31

32 The current study reveals the novel finding that the structure of brain regions in healthy adults
33 varies according to individual abilities in analogical reasoning. These findings highlight
34 structure-function relationships based on individual variations within the normal range in non-
35 pathological subjects. First, a whole-brain VBM analysis showed a negative correlation between
36 GM volume within the left rLPFC and performance on Analogy tasks. VBM-based cluster
37 analyses and SVC analyses using independent ROIs built from a published study on analogical
38 reasoning demonstrated that the left rLPFC was associated with both exemplar-based and
39 concept-based analogy conditions (AnalogyFind and AnalogyApply). This result argues for a
40 role of the left rLPFC in the analogical processes shared by these conditions but not in the
41 perceptual matching task.
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These findings show a strong anatomical convergence with functional imaging. This convergence is of greatest relevance because VBM and functional imaging methods differ in their physiological implications; VBM explores the relationship between brain structure and analogy as variable inter-individual features, while most functional imaging studies that have been performed in this field showed brain regions recruited by all subjects for common analogy processing. These results thus suggest that the rIPFC supports cognitive processes engaged in analogical reasoning and, moreover, that the efficiency of these processes depends on rIPFC morphometry.

In addition to this main result, the whole-brain VBM analysis also pointed to anterior temporal regions, bilateral inferolateral areas being associated with Analogy tasks, while right inferomedial areas were associated with control Match tasks. Tractography showed that these distinct Analogy and Match temporal regions have distinct sets of connections to other regions. These regions may support the distinct semantic and visual processing of information required by Analogy and Match tasks, respectively, as suggested previously (Pascual et al. 2013). Tractography also suggested that the frontal and temporal regions associated with analogy capacity have a different set of connections but share the long segment of the AF that connects both of them.

Altogether, our results suggest that the morphometry of the left rIPFC and anterolateral temporal regions predicts relational reasoning or conceptual abilities rather than perceptual comparisons or similarity identification. As illustrated in Figure S1, perceptual similarities between the source and the target in the Analogy tasks were reduced by varying stimulus attributes such as font, color, position, size, identity, and by introducing cross-dimension analogies in half of the trials. A debriefing questionnaire performed in a previous study that used

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3 the same tasks (Volle et al. 2010) showed that participants were able to verbalize the analogy
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5 schemata used in the tasks at the end of the procedure, suggesting that they processed the
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7 analogies conceptually rather than perceptually.
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11 Finally, this study dissociates the left rIPFC region associated with the mean Analogy
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13 performance (IPOL) from the more posterior lateral PFC region associated with the mean Find
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15 performance (lmidPFC). This result suggests that the left rIPFC (IPOL) supports the relational
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17 and abstract thinking abilities required in analogical reasoning (but not in attribute matching
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19 task), while the left IFG (lmidPFC) is associated with the ability to infer or identify a cognitive
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21 rule based on a perceptual (matching rule) or a relational (analogy schema) similarity. The
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23 tractography analysis showed that the IPOL and lmidPFC regions had different sets of
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25 connections, with IPOL connecting various distant regions while lmidPFC had intra-frontal
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27 connections, with IPOL connecting various distant regions while lmidPFC had intra-frontal
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29 connections.
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33 The role of each of these regions in analogy and the possible physiological meaning of
34
35 these new results will be discussed below.
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41 ***Role of the left rIPFC in analogy***

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44 The current findings suggest that the participants' ability in analogical reasoning depends on
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46 individual variations in GM volume within the left rIPFC. This result converges with previous
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48 findings on the cerebral correlates of analogy using functional imaging (Wharton et al. 2000;
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50 Luo et al. 2003; Christoff et al. 2003; Bunge et al. 2005, 2009; Geake and Hansen 2005, 2010;
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52 Green et al. 2006, 2010; Wendelken et al. 2008, 2012; Wartenburger et al. 2009; Volle et al.
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54 2010; Cho et al. 2010; Krawczyk et al. 2010a; Preusse et al. 2011; Hampshire et al. 2011;
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3 Vartanian 2012; Watson and Chatterjee 2012). Strikingly, the very same left rIPFC region was
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5 observed in functional MRI using similar analogy tasks in different participants (Volle et al.
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8 2010; Figure S2). The experimental design and the results of this study suggest a role for the
9
10 rIPFC in building a structured mental representation of stimulus sets by considering multiple
11
12 relationships. The current findings from the SVC analysis (Table 4) in fact suggest that the left
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14 rIPFC has a role in analogy even when the relational concept is explicitly given (AnalogyApply)
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16 and thus there is no need to find or infer the schema, i.e., no need for concept induction. Both
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18 Analogy conditions still require the processing of relationships between stimuli according to the
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20 task context framed from exemplars or from a verbal term. Therefore, the left rIPFC may be
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22 more involved in considering multiple relationships than in concept induction. This interpretation
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24 is supported by previous fMRI results (Wendelken et al., 2008) showing the involvement of the
25
26 rIPFC in semantic analogy tasks both when participants were to retrieve the relationships
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28 between pairs of words (such as “painter : brush”) and when a relational term (such as “uses”)
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30 was explicitly given. Reinforcing this hypothesis, according to patient studies, prefrontal damage
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32 may cause an impairment in analyzing multiple relationships similar to those employed here
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34 (Krawczyk et al. 2008). Previous studies have demonstrated that the left rIPFC is involved in
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36 analogy tasks whatever the nature of stimuli used (i.e. visuospatial, verbal or semantic), or the
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38 type of analogy schema (mathematical, logical, geometrical, or semantic relation) (Vartanian et
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40 al. 2012). The current results reinforce this idea as the use of various stimuli and heterogeneous
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42 analogy schemata in our Analogy tasks suggests that the left rIPFC involvement in analogy is not
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44 schema-specific. Thus, the current findings add to the existing evidence for a role of the left
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46 rIPFC in domain-general relational integration (Christoff et al., 2001; Reynolds et al. 2006) and
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3 more importantly demonstrate that inter-individual variability in relational reasoning can be
4 supported by structural variation within the left rIPFC.
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9 The rIPFC has been associated with a variety of other cognitive functions and complex
10 cognitive abilities (for reviews see Dumontheil et al. 2008; Ramnani and Owen, 2004) that
11 require the integration of distinct elements of information, such as coordinating goals with
12 subgoals and multitasking (Burgess et al. 2007; Roca et al. 2011), switching attention to
13 stimulus-oriented or stimulus-independent thoughts (Raichle et al. 2001; Gilbert et al. 2005;
14 Burgess et al. 2007), creativity tasks (Gonen-Yaacovi et al. 2013) and fluid intelligence (Geake
15 and Hansen 2005). Whether the rIPFC supports common relational integration processes required
16 by these diverse functions or whether distinct rIPFC subregions have distinct roles in these
17 processes remains to be determined. Exploring the anatomical connections of the rIPFC, an
18 integrative region, may shed some light on this question.
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33 Tractography performed in the current study revealed that the left rIPFC is connected
34 with the semantic system, the ventral visual stream and language areas (temporal and posterior
35 parietal regions) via the AF, IFOF, and UF. It is also connected with the contralateral rostral PFC
36 via callosum fibers and with the thalamus via the anterior thalamic radiations. These connections
37 may enable the integration of the relations among perceived stimuli and the conceptual schema
38 that is either inferred from exemplars (in the AnalogyFind condition) or verbally processed based
39 on the relational term displayed (AnalogyApply).
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52 *Role of temporal areas in analogy*

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3 The current VBM study also revealed a bilateral anterior and inferolateral temporal
4 region for analogy that has not been reported in previous functional MRI studies but has been
5 related to analogical reasoning deficits in patients with temporal damage (Morrison et al. 2004).
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7 In functional MRI, the anterior temporal lobe (ATL) has been rarely reported in association with
8 relational reasoning, partly because this region is often not acquired with MRI or is too sensitive
9 to distortion artifacts caused by magnetic susceptibility, as has been suggested by semantic
10 memory studies (Visser et al. 2010). When reported, the ATL was associated with semantic
11 distance between analogs (Green et al., 2010) or with lower activity during relational reasoning
12 compared with control tasks (Wendelken et al. 2008; Geake and Hansen 2010; Volle et al. 2010).
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14 With VBM methods, the ATL region has been found to correlate with relational reasoning
15 measured by the Raven matrices test (Yuan et al. 2012), which is consistent with our analogy
16 results.
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33 The bilateral ATL is thought to support semantic and/or abstract representations (Hodges
34 et al. 1992; Rogers et al. 2006; Hodges and Patterson 2007; Gorno-Tempini et al 2011) and has
35 been proposed to serve as an amodal (or transmodal) and domain-general semantic “hub,”
36 linking different aspects of knowledge distributed in other brain regions (Patterson et al. 2007;
37 Jefferies 2013). This hypothesis is supported by structural imaging studies in patients with
38 temporal damage (Lambon Ralph et al. 2010, 2012), functional imaging investigations (Vigneau
39 et al. 2006; Binder et al. 2009; Binney et al. 2010; Visser et al. 2010), morphometry (de
40 Zubicaray et al. 2011) and transcranial magnetic stimulation (TMS) studies (Pobric et al. 2009,
41 2010), all showing a critical role of the ATL in category-general semantic tasks. Some of these
42 results especially involved inferolateral ATL subregions very close to our analogy region (Figure
43 S3). Although our Analogy task used symbols, it is possible that analogical reasoning is
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3 fundamentally embedded in the semantic network. This would suggest that concepts stored in the
4 semantic network are necessary to understand any type of analogy, even between symbols, as
5 suggested previously (Knowlton et al. 2012) and may even be verbally formulated (see Volle et
6 al. 2010). Our analogy tasks used various conceptual relations or schemas, which is consistent
7 with the role of the ATL region in semantic memory and in abstracting away from surface
8 similarities (Patterson et al. 2007; Pobric et al. 2010). It is also possible that the left and right
9 ATL reflect verbally and perceptually encoded conceptual representations respectively as our
10 tasks did not allow this distinction (Acres et al. 2009; Gainotti 2012; 2014; Mesulam et al. 2013;
11 Gil-Robles et al. 2013).

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26 The distinct brain correlates of our Analogy and Match tasks suggest a functional
27 dissociation between their neural correlates, with abstract analogies being associated with an
28 inferolateral portion bilaterally and visual similarity with a posterior region and a more medial
29 portion of the right ATL in the fusiform gyrus. The right dominance of the brain correlates of the
30 visual matching task is consistent with the hypothesis of a right inferotemporal advantage for the
31 processing of visual information, while the left temporal cortex may process verbal information
32 (Coello et al. 2013; Gainotti 2014). The fusiform gyrus is part of the ventral stream of visual
33 information processing and allows for the identification of items (Tyler et al. 2013). Visual
34 features had to be matched based on similarity in the Match tasks, which could explain the
35 involvement of this region in the Match tasks.

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50 In the right ATL region, brain correlates of Analogy tasks were more lateral to those of
51 the Match tasks. Anatomical-functional distinctions have been described in the left ATL region
52 (Pascual et al. 2013; Mesulam et al. 2013; Gil-Robles et al. 2013) wherein the inferolateral part
53 is connected to the semantic and default network functionally (Pascual et al. 2013) and
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3 anatomically (Fan et al. 2013) and a more ventral and medial region within the fusiform gyrus is
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5 connected to visual networks. Consistent with this functional specialization, our tractography
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7 results showed a distinct anatomical connectivity between the inferomedial and the inferolateral
8
9 right ATL in addition to their functional orientation toward Analogy and Match performance.
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11 One important difference between the anatomical connectivity patterns of the temporal regions
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13 associated with Analogy and Match performance was that only the inferolateral analogy regions
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15 (ITEMP and rTEMP) were connected to the AF.
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21 The tractography results also showed that the rostral frontal and temporal Analogy
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23 regions shared the AF, which connected the left rIPFC (IPOL) and the left inferolateral ATL
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25 (ITEMP) regions. AF projections extend beyond the classical Broca-Wernicke model (Thiebaut
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27 de Schotten et al. 2012), and AF functions have recently been extended beyond classical
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29 language models including verbal working memory or the ability to learn new “words” (Catani et
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31 al. 2007; Dick and Tremblay 2012; López-Barroso et al. 2013). It is therefore likely that rIPFC
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33 and ATL regions exchange information about the relational concept evoked in Analogy tasks,
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35 although semantic memory is classically associated with UF or IFOF (Duffau et al. 2005, 2008;
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37 de Zubicaray et al. 2011). Finally, we cannot exclude that the rIPFC and ATL regions may also
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39 be indirectly connected anatomically or that these connections are not involved in analogical
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41 reasoning. Overall, the role of AF in analogical reasoning and how the rIPFC and ATL regions
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43 are part of a functional network subserving analogies (de Zubicaray et al. 2011; Wei et al. 2012;
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45 Fan et al. 2013; Pascual et al. 2013) will need further specific exploration.
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55 ***Rule induction is related to a more posterior PFC region***
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3 Our findings also suggest that separate analogy components are associated with distinct lateral
4 prefrontal regions. Whereas reasoning based on relationships rather than on the stimuli
5 themselves is associated with the left rLPFC (IPOL), finding the matching rule or schema is
6 related to the morphology of a more posterior prefrontal region (lmidPFC), whether the rule is
7 concrete (same features) or more abstract (in analogy). This later result is consistent with the role
8 of the lateral left PFC in inductive reasoning demonstrated by lesion studies (Reverberi et al.
9 2005a; 2005b) and functional neuroimaging (Goel and Dolan 2004; Crescentini et al. 2001; Jia et
10 al. 2011; Liang et al. 2014). Studies contrasting rule identification (find the rule) and rule
11 following (apply a given rule) have reported activity in a left lateral prefrontal area very close to
12 our lmidPFC region (Crescentini et al. 2001; Jia et al. 2011). The role of this region in inductive
13 reasoning has been related to its role in detecting regularities across stimuli and generating
14 hypotheses from them (Crescentini et al. 2011). Our paradigm did not allow to clarify the precise
15 cognitive and executive operations supported by the midPFC and engaged in our tasks, and more
16 generally in rule induction. Patient studies also have shown that the left lateral prefrontal cortex
17 is critical for rule finding more than for rule following (Reverberi et al. 2005a, b), suggesting that
18 this region is especially important for inductive reasoning and that the consequences of its
19 damage should be assessed in clinical practice.

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22 Taken together, these findings suggest that distinct brain regions support distinct analogy
23 processes: the left rLPFC may support the processing and integration of the relationships between
24 stimuli enabling the representation of a conceptual schema in interaction with the ATL regions
25 involved in semantic memory. A more posterior lateral PFC region may be engaged in the
26 inference processes required to identify a matching rule in exemplar-based conditions, whether
27 the rule is concrete (a perceptual match) or more abstract (an analogy schema). The two distinct
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3 prefrontal regions we observed may also be linked to the recent models describing a caudal-
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5 rostral organization of prefrontal cortex according to distinct levels and types of abstraction
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7 (Badre 2008 for a review), in which most anterior prefrontal regions support more abstract rules
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9 or action representations or complex relational/semantic integration (Green et al. 2006) than
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11 more posterior ones.
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22 *Interpretation of correlations with brain structures*

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24 We observed negative correlations between local GM volume and performance in both the
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26 Analogy and Match conditions. In other non-frontal regions, we also observed positive
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28 correlations. In both cases, these regions were found to be activated in functional imaging studies
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30 using related tasks or processes. The common notion of “the more brain volume the better”
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32 appears to need reconsideration, at least in some cerebral regions. While some of the evidence
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34 does suggest that greater local GM volume is associated with better performance, especially
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36 when comparing patients to controls or after a specific training or acquired expertise (Maguire et
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38 al. 2000; Draganski and May 2008; Takeuchi et al. 2010, 2011, 2013), others report better
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40 performance associated with less local GM volume, especially in the prefrontal regions (Moore
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42 et al. 2009; Buda et al. 2011; Goh et al. 2011; Smolker et al. 2014). Several studies also showed
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44 both positive and negative correlations for different brain regions and different aspects of multi-
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46 faceted concepts, such as empathy (Banissy et al. 2012), intelligence (Frangou et al. 2004), or
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48 creativity (Jung et al., 2010). Overall, it appears that the observation of positive or negative
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50 structure-task correlations depends heavily on brain regions, on the population studied, and on
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52 the particular process assessed. Thus, it is important to consider why less GM signal in VBM
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3 would actually facilitate performance. A first possibility may be that less GM volume allows
4 greater WM volume (i.e., less cell bodies and more connections; Paus 2005; Goh et al. 2011).
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6 This may be especially relevant for rostral PFC, where dendritic neuropil is more developed than
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8 in other comparable areas of the cortex while the density of cell bodies is low, pointing to its
9
10 high integrative properties (Ramnani and Owen 2004; Dumontheil et al. 2008). Alternatively, the
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12 process of synaptic pruning that occurs during brain maturation and leads to frontal cortex
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14 thinning (Dumontheil et al. 2008; Shaw et al. 2006) has also been associated with an
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16 improvement in executive functions (Kharitonova and Munakata 2011). Cortical thickness has
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18 been specifically measured in an analogy study on adolescents (Krawczyk et al., 2010b) that
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20 demonstrated a correlation between rIPFC thinning and a better performance in analogical
21
22 reasoning. This study pointed to a more medial rIPFC region than the current results, likely be
23
24 due to a difference in the material used, i.e. meaningful pictures with distinct difficulty levels.
25
26 These findings reinforce the hypothesis that brain development that leads to a thinner left rIPFC
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28 may confer higher analogical abilities. Differences in local brain development may also explain
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30 variations observed in the left rIPFC involvement in analogy tasks during childhood (Dumontheil
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32 et al. 2010). In this context, although the significance of “macroscopic” anatomical variations is
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34 not yet understood in terms of microscopic variability (such as cellular or synaptic variability),
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36 possible interpretations of our data are that good performers may have experienced a more
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38 efficient synaptic pruning or cortical myelination, leading to a thinner rostral PFC cortex.
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49 Our findings show that the macroscopic correlates of cognitive abilities are not
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51 homogeneous across regions, suggesting that the mechanisms supporting the cognitive capacities
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53 of the cortex may be distinct in the frontal and temporal regions. These mechanisms may rely
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55 differently on cellular or neuropil changes, or on surface folding or cortical thickening (Mechelli
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3 et al. 2005), and may be related to both genetic and environmental factors. A better
4
5 understanding of the physiological bases of local increases and decreases in GM volume will be
6
7 necessary to interpret our VBM results (Kanai and Rees 2011; Eriksson et al. 2009). It is
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9 noteworthy that in functional MRI, the biological significance of activation and deactivation is
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11 not fully understood either (Logothetis 2008).
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15 As the performances in our Analogy and control Match tasks were not equivalent, we can
16
17 not exclude that the difference in their cerebral correlates could be related to difficulty-related
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19 processing. However, we believe that the brain correlates of our Analogy tasks reflect analogical
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21 reasoning abilities, because our findings are consistent with previous studies that controlled
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23 difficulty levels (Hampshire et al. 2011; Kroger et al. 2002; Cho et al. 2010) or equated
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25 performance between analogy and control tasks (Watson et al. 2012; Krawczyk et al. 2010a).
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27 Activation within the same rIPFC region has been consistently reported during tasks that involve
28
29 analogical reasoning whatever the difficulty or perceptual nature of the material used (see a
30
31 meta-analysis from Vartanian et al. 2012). Furthermore, in a previous study (Wartenburger et al.
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33 2009) the left rIPFC was only moderately modulated by difficulty in analogical reasoning.
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40 Finally, despite the large age range of the participants, it is unlikely that aging may have
41
42 biased the results for several reasons. First, our analyses were corrected for age and total GM
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44 volume. Second, the correlation between GM volume and age within the left rIPFC region
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46 (IPOL) disappeared when controlling for the total GM volume, which does not argue for a local
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48 effect of ageing on the GM volume of the rIPFC.
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55 **Conclusion**

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The originality of this study was to investigate the structural correlates of analogical reasoning using VBM coupled with an anatomical connectivity study. Combined with previous functional imaging reports, our results suggest that the left rIPFC and inferolateral ATL support cognitive processes engaged when solving analogies and that variability in their anatomy predicts individual differences in the efficiency of this processing. Considering previous reports, the profile of the brain correlates observed in our distinct experimental conditions suggests that variability in the left rIPFC structure and the temporal semantic regions may reflect an ability to process multiple relationships between stimuli and to link them to a conceptual schema. The ability to infer the analogy schema from exemplars may depend on a more posterior left lateral PFC region. Further research would be necessary to deepen our understanding of the roles these regions play in reasoning processes and their relationship to recent models of prefrontal organization, to clarify how they interact via their anatomical or functional connectivity, and to examine how damage to these areas or their connections provokes an impairment in analogy abilities. Other methods that allow stronger causality, such as lesion studies or TMS, would allow for drawing more definitive conclusions about the involvement of the left rIPFC in reasoning. The very few existing pioneering studies in this direction indeed highlighted the importance of an intact PFC for relational reasoning (Boroojerdi et al. 2001; Morrison et al. 2004; Krawczyk et al. 2008; Schmidt et al. 2012). The current findings offer new anatomical targets for future research on the cognitive consequences of damage to these regions and their connections.

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For Peer Review

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For Peer Review

References

- Acres K, Taylor KI, Moss, HE, Stamatakis EA, Tyler LK. 2009. Complementary hemispheric asymmetries in object naming and recognition: A voxel-based correlational study. *Neuropsychologia*. 47: 1836-1843. doi:10.1016/j.neuropsychologia.2009.02.024
- Alexander DC. 2006. An Introduction to Computational Diffusion MRI: the Diffusion Tensor and Beyond. In: Weickert PJ, Hagen PH, editors. *Visualization and Processing of Tensor Fields*. Springer Berlin Heidelberg. p. 83–106.
- Anderson AW. 2005. Measurement of fiber orientation distributions using high angular resolution diffusion imaging. *Magn Reson Med*. 54:1194–1206.
- Ashburner J. 2007. A fast diffeomorphic image registration algorithm. *Neuroimage*. 38:95–113.
- Ashburner J, Friston KJ. 2000. Voxel-based morphometry--the methods. *Neuroimage*. 11:805–821.
- Ashburner J, Friston KJ. 2005. Unified segmentation. *Neuroimage*. 26:839–851.
- Badre D, Wagner AD. 2007. Left ventrolateral prefrontal cortex and the cognitive control of memory. *Neuropsychologia*. 45:2883–2901.
- Badre, D. 2008. Cognitive control, hierarchy, and the rostro-caudal organization of the frontal lobes. *Trends Cogn Sci*. 12:193-200.
- Banissy MJ, Kanai R, Walsh V, Rees G. 2012. Inter-individual differences in empathy are reflected in human brain structure. *Neuroimage*. 62:2034–2039.

- 1
2
3 Binder JR, Desai RH, Graves WW, Conant LL. 2009. Where Is the Semantic System? A Critical
4
5 Review and Meta-Analysis of 120 Functional Neuroimaging Studies. *Cereb Cortex*.
6
7 19:2767–2796.
8
9
- 10
11 Binney RJ, Embleton K V, Jefferies E, Parker GJM, Lambon Ralph MA. 2010. The ventral and
12
13 inferolateral aspects of the anterior temporal lobe are crucial in semantic memory: evidence
14
15 from a novel direct comparison of distortion-corrected fMRI, rTMS, and semantic
16
17 dementia. *Cereb Cortex*. 20:2728–2738.
18
19
- 20
21
22 Blanchette I, Dunbar K. 2000. How analogies are generated: the roles of structural and
23
24 superficial similarity. *Mem Cogn*. 28:108–124.
25
26
27
- 28
29 Boroojerdi B, Phipps M, Kopylev L, Wharton CM, Cohen LG, Grafman J. 2001. Enhancing
30
31 analogic reasoning with rTMS over the left prefrontal cortex. *Neurology*. 56:526–528.
32
33
- 34
35 Buda M, Fornito A, Bergström ZM, Simons JS. 2011. A Specific Brain Structural Basis for
36
37 Individual Differences in Reality Monitoring. *J Neurosci*. 31:14308–14313.
38
39
- 40
41 Bunge SA, Helskog EH, Wendelken C. 2009. Left, but not right, rostrolateral prefrontal cortex
42
43 meets a stringent test of the relational integration hypothesis. *Neuroimage*. 46:338–342.
44
45
- 46
47 Bunge SA, Wendelken C, Badre D, Wagner AD. 2005. Analogical reasoning and prefrontal
48
49 cortex: evidence for separable retrieval and integration mechanisms. *Cereb Cortex*. 15:239–
50
51 249.
52
53
- 54
55 Burgess PW, Gilbert SJ, Dumontheil I. 2007. Function and localization within rostral prefrontal
56
57 cortex (area 10). *Philos Trans R Soc London - Ser B Biol Sci*. 362:887–899.
58
59
60

- 1
2
3 Catani M, Allin MPG, Husain M, Pugliese L, Mesulam MM, Murray RM, Jones DK. 2007.
4
5 Symmetries in human brain language pathways correlate with verbal recall. *Proc Natl Acad*
6
7
8 *Sci U S A.* 104:17163–17168.
9
10
11 Catani M, Dell’Acqua F, Vergani F, Malik F, Hodge H, Roy P, Valabregue R, Thiebaut de
12
13 Schotten M. 2012. Short frontal lobe connections of the human brain. *Cortex.* 48:273–291.
14
15
16
17 Cho S, Moody TD, Fernandino L, Mumford J a, Poldrack R a, Cannon TD, Knowlton BJ,
18
19 Holyoak KJ. 2010. Common and dissociable prefrontal loci associated with component
20
21 mechanisms of analogical reasoning. *Cereb Cortex.* 20:524–533.
22
23
24
25
26 Christoff K, Keramatian K, Gordon AM, Smith R, Mädler B. 2009. Prefrontal organization of
27
28 cognitive control according to levels of abstraction. *Brain Res.* 1286:94–105.
29
30
31
32 Christoff K, Prabhakaran V, Dorfman J, Zhao Z, Kroger JK, Holyoak KJ, Gabrieli JD. 2001.
33
34 Rostrolateral prefrontal cortex involvement in relational integration during reasoning.
35
36 *Neuroimage.* 14:1136–1149.
37
38
39
40 Christoff K, Ream JM, Geddes LPT, Gabrieli JDE. 2003. Evaluating self-generated information:
41
42 anterior prefrontal contributions to human cognition. *Behav Neurosci.* 117:1161–1168.
43
44
45
46 Coello, A. F., Duvaux, S., De Benedictis, A., Matsuda, R., & Duffau, H. 2013. Involvement of
47
48 the right inferior longitudinal fascicle in visual hemiagnosia: a brain stimulation mapping
49
50 study: Case report. *Journal of neurosurgery.* 118: 202–205.
51
52
53
54
55
56
57
58
59
60

1
2
3 Colom R, Haier RJ, Head K, Álvarez-linera J, Ángeles M, Chun P, Jung RE. 2009. Intelligence
4
5 Gray matter correlates of fluid, crystallized, and spatial intelligence: Testing the P-FIT
6
7 model. *Intelligence*. 37:124–135.
8
9

10
11 Colom R, Jung RE, Haier RJ. 2006. Distributed brain sites for the g-factor of intelligence.
12
13 *Neuroimage*. 31:1359–1365.
14
15

16
17 Crescentini C, Seyed-Allaei S, De Pisapia N, Jovicich J, Amati D, Shallice T. 2011. Mechanisms
18
19 of Rule Acquisition and Rule Following in Inductive Reasoning. *The Journal of*
20
21 *Neuroscience*, 31: 7763 -7774. doi:10.1523/JNEUROSCI.4579-10.2011
22
23

24
25 Crone EA, Wendelken C, van Leijenhorst L, Honomichl RD, Christoff K, Bunge SA. 2009.
26
27 Neurocognitive development of relational reasoning. *Dev Sci*. 12:55–66.
28
29

30
31 De Zubicaray GI, Rose SE, McMahon KL. 2011. The structure and connectivity of semantic
32
33 memory in the healthy older adult brain. *Neuroimage*. 54:1488–1494.
34
35

36
37 Dell'Acqua F, Scifo P, Rizzo G, Catani M, Simmons A, Scotti G, Fazio F. 2010. A modified
38
39 damped Richardson-Lucy algorithm to reduce isotropic background effects in spherical
40
41 deconvolution. *Neuroimage*. 49:1446–1458.
42
43

44
45 Dell'Acqua F, Simmons A, Williams SCR, Catani M. 2013. Can spherical deconvolution
46
47 provide more information than fiber orientations? Hindrance modulated orientational
48
49 anisotropy, a true-tract specific index to characterize white matter diffusion. *Hum Brain*
50
51 *Mapp*. 34:2464–2483.
52
53
54
55
56
57
58
59
60

- 1
2
3 Dick AS, Tremblay P. 2012. Beyond the arcuate fasciculus: consensus and controversy in the
4
5 connectional anatomy of language. *Brain*. 135:3529–3550.
6
7
8
9 Draganski B, May A. 2008. Training-induced structural changes in the adult human brain. *Behav*
10
11 *Brain Res*. 192:137–142.
12
13
14
15 Dubois B, Slachevsky A, Litvan I, Pillon B. 2000. The FAB: a Frontal Assessment Battery at
16
17 bedside. *Neurology*. 57:1621–1626.
18
19
20
21 Duffau H, Gatignol P, Mandonnet E, Peruzzi P, Tzourio-Mazoyer N, Capelle L. 2005. New
22
23 insights into the anatomo-functional connectivity of the semantic system: a study using
24
25 cortico-subcortical electrostimulations. *Brain*. 128:797–810.
26
27
28
29 Duffau H, Peggy Gatignol ST, Mandonnet E, Capelle L, Taillandier L. 2008. Intraoperative
30
31 subcortical stimulation mapping of language pathways in a consecutive series of 115
32
33 patients with Grade II glioma in the left dominant hemisphere. *J Neurosurg*. 109:461–471.
34
35
36
37 Dumontheil I, Burgess PW, Blakemore S-J. 2008. Development of rostral prefrontal cortex and
38
39 cognitive and behavioural disorders. *Dev Med Child Neurol*. 50:168–181.
40
41
42
43 Dumontheil I, Houlton R, Christoff K, Blakemore S-J. 2010. Development of relational
44
45 reasoning during adolescence. *Dev Sci*. 13:F15–24.
46
47
48
49 Einstein A. 1938. On the Effects of External Sensory Input on Time Dilation. *J Exothermic Sci*
50
51 *Technol*. 1(9).
52
53
54
55
56
57
58
59
60

- 1
2
3 Eriksson SH, Free SL, Thom M, Symms MR, Martinian L, Duncan JS, Sisodiya SM. 2009.
4
5 Quantitative grey matter histological measures do not correlate with grey matter probability
6
7 values from in vivo MRI in the temporal lobe. *Journal of Neuroscience Methods*. 181:
8
9 111-118.
10
11
12
13
14 Fan L, Wang J, Zhang Y, Han W, Yu C, Jiang T. 2013. Connectivity-Based Parcellation of the
15
16 Human Temporal Pole Using Diffusion Tensor Imaging. *Cereb Cortex*. 2013 Aug 7. [Epub
17
18 ahead of print]
19
20
21
22
23 Folstein MF, Folstein SE, McHugh PR. 1975. "Mini-Mental State" - A practical method for
24
25 grading the cognitive state of patients for the clinician. *J Psychisychiatric Res*. 12:189-198.
26
27
28
29 Frangou S, Chitins X, Williams SCR. 2004. Mapping IQ and gray matter density in healthy
30
31 young people. *Neuroimage*. 23:800-805.
32
33
34
35 Gainotti G. 2012. The format of conceptual representations disrupted in semantic dementia: A
36
37 position paper. *Cortex*. 48: 521-529. doi:10.1016/j.cortex.2011.06.019
38
39
40
41 Gainotti G. 2014. Why Are the Right and Left Hemisphere Conceptual Representations
42
43 Different? *Behavioural Neurology*. 603134. doi:10.1155/2014/603134
44
45
46
47 Geake JG, Hansen PC. 2005. Neural correlates of intelligence as revealed by fMRI of fluid
48
49 analogies. *Neuroimage*. 26:555-564.
50
51
52
53 Geake JG, Hansen PC. 2010. Functional neural correlates of fluid and crystallized analogizing.
54
55
56
57
58
59
60

1
2
3 Gentner D. 1983. Structure-Mapping: A Theoretical Framework for Analogy. *Cogn Sci.* 7:155–
4
5 170.

6
7
8
9 Gentner D, Holyoak KJ. 1997. Reasoning and learning by analogy. *Am Psychol.* 52:32–34.

10
11
12 Gentner D, Medina J. 1998. Similarity and the development of rules. *Cognition.* 65:263–297.

13
14
15
16 Gilbert SJ, Frith CD, Burgess PW. 2005. Involvement of rostral prefrontal cortex in selection
17
18 between stimulus-oriented and stimulus-independent thought. *Eur J Neurosci.* 21:1423–
19
20 1431.

21
22
23
24
25 Gil-Robles S, Carvallo A, Jimenez M, Gomez Caicoya A, Martinez R, Ruiz-Ocaña, C, Duffau H.
26
27 2013. Double dissociation between visual recognition and picture naming: a study of the
28
29 visual language connectivity using tractography and brain stimulation. *Neurosurgery.* 72:
30
31 678-686. doi:10.1227/NEU.0b013e318282a361

32
33
34
35
36 Goel V and Dolan RJ. 2004. Differential involvement of left prefrontal cortex in inductive and
37
38 deductive reasoning. *Cognition.* 93: B109-B121. doi:10.1016/j.cognition.2004.03.001

39
40
41 Goh S, Bansal R, Xu D, Hao X, Liu J, Peterson BS. 2011. Neuroanatomical correlates of
42
43 intellectual ability across the life span. *Dev Cogn Neurosci.* 1:305–312.

44
45
46
47 Gonen-Yaacovi G, de Souza LC, Levy R, Urbanski M, Josse G, Volle E. 2013. Rostral and
48
49 caudal prefrontal contribution to creativity: a meta-analysis of functional imaging data.
50
51 *Front Hum Neurosci.* 7:465.

- 1
2
3 Good CD, Johnsrude IS, Ashburner J, Henson RNA, Friston KJ, Frackowiak RSJ. 2001. A
4 voxel-based morphometric study of ageing in 465 normal adult human brains. *Neuroimage*.
5 14:21–36.
6
7
8
9
10
11 Gorno-Tempini ML, Hillis AE, Weintraub S, Kertesz A, Mendez M, Cappa SF, Ogar JM, Rohrer
12 JD, Black S, Boeve BF, Manes F, Dronkers NF, Vandenberghe R, Rascovsky K, Patterson
13 K, Miller BL, Knopman DS, Hodges JR, Mesulam MM, Grossman M. 2011. Classification
14 of primary progressive aphasia and its variants. *Neurology*. 76:1006–1014.
15
16
17
18
19
20
21
22 Green AE, Fugelsang J a, Kraemer DJM, Shamosh N a, Dunbar KN. 2006. Frontopolar cortex
23 mediates abstract integration in analogy. *Brain Res*. 1096:125–137.
24
25
26
27
28 Green AE, Kraemer DJM, Fugelsang J a, Gray JR, Dunbar KN. 2010. Connecting long distance:
29 semantic distance in analogical reasoning modulates frontopolar cortex activity. *Cereb*
30 *Cortex*. 20:70–76.
31
32
33
34
35
36 Green AE, Kraemer DJM, Fugelsang J a, Gray JR, Dunbar KN. 2012. Neural correlates of
37 creativity in analogical reasoning. *J Exp Psychol Learn Mem Cogn*. 38:264–272.
38
39
40
41
42 Haier RJ, Jung RE, Yeo RA, Head K, Alkire MT. 2004. Structural brain variation and general
43 intelligence. *Neuroimage*. 23:425–433.
44
45
46
47
48 Hampshire A, Thompson R, Duncan J, Owen AM. 2011. Lateral prefrontal cortex subregions
49 make dissociable contributions during fluid reasoning. *Cereb Cortex*. 21:1–10.
50
51
52
53
54 Jung RE and Haier RJ. 2007. The Parieto-Frontal Integration Theory (P-FIT) of intelligence:
55 converging neuroimaging evidence. *Behav Brain Sci*. 30: 135-54; discussion 154-87.
56
57
58
59
60

1
2
3 Knowlton BJ, Morrison RG, Hummel JE, Holyoak KJ. 2012. A neurocomputational system for
4 relational reasoning. *Trends in Cognitive Sciences*. 16: 373–381.
5
6

7
8
9 Hodge, JR and Patterson, K. 2007. Semantic dementia: a unique clinicopathological syndrome.
10
11 *The Lancet Neurology*. 6: 1004-1014. doi:10.1016/S1474-4422(07)70266-1
12
13

14
15 Hodges JR, Patterson K, Oxbury S., Funnell E. 1992. Semantic dementia. Progressive fluent
16
17 aphasia with temporal lobe atrophy. *Brain*. 115 (Pt 6): 1783-1806.
18
19

20
21 Hofstadter D, Sander E. 2013. *Surfaces and Essences: Analogy as the Fuel and Fire of Thinking*.
22
23 New York: Basic Books.
24
25

26
27 Holyoak KJ, Morrison RG. 2012. Analogy and Relational Reasoning. In: Holyoak KJ, Morrison,
28
29 editors. *The Oxford handbook of thinking and reasoning*. New York: Oxford University
30
31 Press. p. 234–259.
32
33

34
35 Holyoak KJ, Thagard P. 1995. *Mental leaps: Analogy in creative thought*. Cambridge, MA: MIT
36
37 Press.
38
39

40
41 Jefferies E. 2013. The neural basis of semantic cognition: Converging evidence from
42
43 neuropsychology, neuroimaging and TMS. *Cortex*. 49:611–625.
44
45

46
47 Jia X, Liang P, Lu J, Yang Y, Zhong N, Li K. 2011. Common and dissociable neural correlates
48
49 associated with component processes of inductive reasoning. *NeuroImage*. 56: 2292-2299.
50
51 doi:10.1016/j.neuroimage.2011.03.020
52
53
54
55
56
57
58
59
60

- 1
2
3 Jung RE, Segall JM, Bockholt HJ, Flores RA, Shirley M, Chavez RS, Haier RJ. 2010.
4
5 Neuroanatomy of Creativity. *Hum Brain Mapp.* 31:398–409.
6
7
8
9 Kanai R, Rees G. 2011. The structural basis of inter-individual differences in human behaviour
10
11 and cognition. *Nat Rev Neurosci.* 12:231–242.
12
13
14
15 Kharitonova M, Munakata Y. 2011. The Role of Representations in Executive Function:
16
17 Investigating a Developmental Link between Flexibility and Abstraction. *Front Psychol.*
18
19 2:347.
20
21
22
23 Klein A, Andersson J, Ardekani BA, Ashburner J, Avants B, Chiang MC, Christensen GE,
24
25 Collins DL, Gee J, Hellier P, Song JH, Jenkinson M, Lepage C, Rueckert D, Thompson P,
26
27 Vercauteren T, Woods RP, Mann JJ, Parsey R V. 2009. Evaluation of 14 nonlinear
28
29 deformation algorithms applied to human brain MRI registration. *Neuroimage.* 46:786–802.
30
31
32
33
34 Koechlin E, Basso G, Pietrini P, Panzer S, Grafman J. 1999. The role of the anterior prefrontal
35
36 cortex in human cognition. *Nature.* 399:148–151.
37
38
39
40 Krawczyk DC. 2012. The cognition and neuroscience of relational reasoning. *Brain Res.*
41
42 1428:13–23.
43
44
45
46 Krawczyk DC, McClelland MM, Donovan CM, Tillman GD, Maguire MJ. 2010a. An fMRI
47
48 investigation of cognitive stages in reasoning by analogy. *Brain Res.* 1342:63–73.
49
50
51
52 Krawczyk DC, Hanten G, Wilde EA, Li X, Schnelle KP, Merkley TL, Vasquez AC, Cook LG,
53
54 McClelland M, Chapman SB, Levin HS. 2010b. Deficits in Analogical Reasoning in
55
56 Adolescents with Traumatic Brain Injury. *Frontiers in human neuroscience.* 4:1-13.
57
58
59
60

- 1
2
3 Krawczyk DC, Morrison RG, Viskontas I, Holyoak KJ, Chow TW, Mendez MF, Miller BL,
4
5 Knowlton BJ. 2008. Distraction during relational reasoning: the role of prefrontal cortex in
6
7 interference control. *Neuropsychologia*. 46:2020–2032.
8
9
10
11 Kroger JK, Sabb FW, Fales CL, Bookheimer SY, Cohen MS, Holyoak KJ. 2002. Recruitment of
12
13 anterior dorsolateral prefrontal cortex in human reasoning: a parametric study of relational
14
15 complexity. *Cereb Cortex*. 12: 477-85.
16
17
18
19
20 Lambon Ralph MA, Cipolotti L, Manes F, Patterson K. 2010. Taking both sides: do unilateral
21
22 anterior temporal lobe lesions disrupt semantic memory? *Brain*. 133:3243–3255.
23
24
25
26 Lambon Ralph MA, Ehsan S, Baker G a, Rogers TT. 2012. Semantic memory is impaired in
27
28 patients with unilateral anterior temporal lobe resection for temporal lobe epilepsy. *Brain*.
29
30 135:242–258.
31
32
33
34 Liang P, Jia X, Taatgen NA, Zhong N, Li K. 2014. Different strategies in solving series
35
36 completion inductive reasoning problems: An fMRI and computational study. *International*
37
38 *Journal of Psychophysiology*. 93:253-260. doi:10.1016/j.ijpsycho.2014.05.006
39
40
41
42 Logothetis NK. 2008. What we can do and what we cannot do with fMRI. *Nature*. 453:869–878.
43
44
45
46 López-Barroso D, Catani M, Ripollés P, Dell’Acqua F, Rodríguez-Fornells A, de Diego-
47
48 Balaguer R. 2013. Word learning is mediated by the left arcuate fasciculus. *Proc Natl Acad*
49
50 *Sci U S A*. 110:13168–13173.
51
52
53
54 Luo Q, Perry C, Peng D, Jin Z, Xu D, Ding G. 2003. The neural substrate of analogical
55
56 reasoning : an fMRI study. *Cog Brain Research*. 17:527–534.
57
58
59
60

- 1
2
3 Maguire EA, Gadian DG, Johnsrude IS, Good CD, Ashburner J, Frackowiak RS, Frith CD. 2000.
4
5 Navigation-related structural change in the hippocampi of taxi drivers. *Proc Natl Acad Sci*
6
7 U S A. 97:4398–4403.
8
9
10
11 Markman AB, Gentner D. 1993. Structural Alignment during Similarity Comparisons. *Cogn*
12
13 *Psychol.* 25:431–467.
14
15
16
17 Markman AB, Gentner D. 2000. Structure mapping in the comparison process. *Am J Psychol.*
18
19 113:501–538.
20
21
22
23 Mechelli A, Price CJ, Friston KJ, Ashburner J. 2005. Voxel-Based Morphometry of the Human
24
25 Brain: Methods and Applications. *Current Medical Imaging Reviews.* 1: 105.
26
27
28
29 Mesulam MM, Wieneke C, Hurley R, Rademaker A, Thompson CK, Weintraub S, Rogalski EJ.
30
31 2013. Words and objects at the tip of the left temporal lobe in primary progressive aphasia.
32
33 *Brain.* 136: 601-618. doi:10.1093/brain/aws336
34
35
36
37
38 Montgomery A, Asberg M. 1979. A New Depression Scale Designed to be Sensitive to Change.
39
40 *Br J psychiatry.* 134:382–389.
41
42
43
44 Moore DW, Bhadelia R a, Billings RL, Fulwiler C, Heilman KM, Rood KMJ, Gansler D a. 2009.
45
46 Hemispheric connectivity and the visual-spatial divergent-thinking component of creativity.
47
48 *Brain Cogn.* 70:267–272.
49
50
51
52 Morrison RG, Krawczyk DC, Holyoak KJ, Hummel JE, Chow TW, Miller BL, Knowlton BJ.
53
54 2004. A neurocomputational model of analogical reasoning and its breakdown in
55
56 frontotemporal lobar degeneration. *J Cogn Neurosci.* 16:260–271.
57
58
59
60

- 1
2
3 Narr KL, Woods RP, Thompson PM, Szeszko P, Robinson D, Dimtcheva T, Gurbani M, Toga
4
5 AW, Bilder RM. 2007. Relationships between IQ and regional cortical gray matter
6
7 thickness in healthy adults. *Cereb Cortex*. 17:2163–2171.
8
9
10
11 Pascual B, Masdeu JC, Hollenbeck M, Makris N, Insausti R, Ding S-L, Dickerson BC. 2013.
12
13 Large-Scale Brain Networks of the Human Left Temporal Pole: A Functional Connectivity
14
15 MRI Study. *Cereb Cortex*. 2013 Sep 24. [Epub ahead of print]
16
17
18
19
20 Patterson K, Nestor PJ, Rogers TT. 2007. Where do you know what you know? The
21
22 representation of semantic knowledge in the human brain. *Nat Rev Neurosci*. 8:976–987.
23
24
25
26 Paus T. 2005. Mapping brain maturation and cognitive development during adolescence. *Trends*
27
28 *Cogn Sci*. 9:60–68.
29
30
31
32 Pobric G, Jefferies E, Lambon Ralph MA. 2010. Category-Specific versus Category-General
33
34 Semantic Impairment Induced by Transcranial Magnetic Stimulation. *Curr Biol*. 20:964–
35
36 968.
37
38
39
40 Pobric G, Lambon Ralph MA, Jefferies E. 2009. The role of the anterior temporal lobes in the
41
42 comprehension of concrete and abstract words: rTMS evidence. *Cortex*. 45:1104–1110.
43
44
45
46 Preusse F, van der Meer Elke, Deshpande G, Krueger F, Wartenburger I. 2011. Fluid intelligence
47
48 allows flexible recruitment of the parieto-frontal network in analogical reasoning. *Front*
49
50 *Hum Neurosci*. 5:22.
51
52
53
54 Raichle ME, MacLeod AM, Snyder AZ, Powers WJ, Gusnard DA, Shulman GL. 2001. A default
55
56 mode of brain function. *Proc Natl Acad Sci U S A*. 98:676–682.
57
58
59
60

- 1
2
3 Ramnani N, Owen AM. 2004. Anterior prefrontal cortex: insights into function from anatomy
4 and neuroimaging. *Nat Rev Neurosci.* 5:184–194.
5
6
7
8
9 Reverberi C, Lavaroni A, Gigli GL, Skrap M, Shallice T. 2005a. Specific impairments of rule
10 induction in different frontal lobe subgroups. *Neuropsychologia.* 43: 460-472.
11
12 doi:10.1016/j.neuropsychologia.2004.06.008
13
14
15
16
17 Reverberi C, D’Agostini, S, Skrap M, Shallice T. 2005b. Generation and recognition of abstract
18 rules in different frontal lobe subgroups. *Neuropsychologia.* 43: 1924-37.
19
20
21
22
23 Reynolds JR, McDermott KB, Braver TS. 2006. A direct comparison of anterior prefrontal
24 cortex involvement in episodic retrieval and integration. *Cereb Cortex.* 16:519–528.
25
26
27
28
29 Ridgway GR, Henley SMD, Rohrer JD, Scahill RI, Warren JD, Fox NC. 2008. Ten simple rules
30 for reporting voxel-based morphometry studies. *Neuroimage.* 40:1429–1435.
31
32
33
34
35 Roca M, Torralva T, Gleichgerrcht E, Woolgar A, Thompson R, Duncan J, Manes F. 2011. The
36 role of Area 10 (BA10) in human multitasking and in social cognition: a lesion study.
37
38 *Neuropsychologia.* 49:3525–3531.
39
40
41
42
43 Rogers TT, Hocking J, Noppeney U, Mechelli A, Gorno-Tempini ML, Patterson K, Price CJ.
44
45 2006. Anterior temporal cortex and semantic memory: reconciling findings from
46
47 neuropsychology and functional imaging. *Cognitive, Affective, & Behavioral Neuroscience.*
48
49 6: 201–213.
50
51
52
53
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46
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52
53
54
55
56
57
58
59
60
- Rojkova K, Volle E, Urbanski M, Humbert F, Dell'Acqua F, Thiebaut de Schotten M. Atlasing the frontal lobe connections and their variability due to age and education: a spherical deconvolution tractography study. *Neuroimage. Under revision.*
- Schmahmann JD, Pandya DN, Wang R, Dai G, D'Arceuil HE, de Crespigny AJ, Wedeen VJ. 2007. Association fibre pathways of the brain: parallel observations from diffusion spectrum imaging and autoradiography. *Brain a J Neurol.* 130:630–653.
- Schmidt GL, Cardillo ER, Kranjec A, Lehet M, Widick P, Chatterjee A. 2012. Not all analogies are created equal: Associative and categorical analogy processing following brain damage. *Neuropsychologia.* 50:1372–1379.
- Shaw P, Greenstein D, Lerch J, Clasen L, Lenroot R, Gogtay N, Evans A, Rapoport J, Giedd J. 2006. Intellectual ability and cortical development in children and adolescents. *Nature.* 440:676-9.
- Smolker HR, Depue BE, Reineberg AE, Orr JM, Banich MT. 2014. Individual differences in regional prefrontal gray matter morphometry and fractional anisotropy are associated with different constructs of executive function. *Brain Struct Funct.* 2014 Feb 22. [Epub ahead of print]
- Takeuchi H, Taki Y, Hashizume H, Sassa Y, Nagase T, Nouchi R, Kawashima R. 2011. Effects of training of processing speed on neural systems. *J Neurosci.* 31:12139–12148.

- 1
2
3 Takeuchi H, Taki Y, Sassa Y, Hashizume H, Sekiguchi A, Fukushima A, Kawashima R. 2010.
4
5 Regional gray matter volume of dopaminergic system associate with creativity: evidence
6
7 from voxel-based morphometry. *Neuroimage*. 51:578–585.
8
9
10
11 Takeuchi H, Taki Y, Nouchi R, Hashizume H, Sekiguchi A, Kotozaki Y, Nakagaw S, Miyauchi
12
13 CM, Sassa Y, Kawashima R. 2013. Effects of multitasking-training on gray matter structure
14
15 and resting state neural mechanisms. *Hum Brain Mapp*. 2013 Dec 17. doi:
16
17 10.1002/hbm.22427. [Epub ahead of print]
18
19
20
21
22 Thibaut JP, French R, Vezneva M. 2010a. The development of analogy making in children:
23
24 Cognitive load and executive functions. *Journal of Experimental Child Psychology*. 106:
25
26 1-19.
27
28
29
30 Thibaut JP, French R, Vezneva M. 2010b. Cognitive load and semantic analogies: Searching
31
32 semantic space. *Psychonomic Bulletin & Review*. 17:569-574.
33
34
35
36 Thiebaut de Schotten M, Dell'Acqua F, Valabregue R, Catani M. 2012. Monkey to human
37
38 comparative anatomy of the frontal lobe association tracts. *Cortex*. 48:82–96.
39
40
41
42 Thiebaut de Schotten M, Ffytche DH, Bizzi A, Dell'Acqua F, Allin M, Walshe M, Murray R,
43
44 Williams SC, Murphy DGM, Catani M. 2011. Atlasing location, asymmetry and inter-
45
46 subject variability of white matter tracts in the human brain with MR diffusion tractography.
47
48 *Neuroimage*. 54:49–59.
49
50
51
52
53
54
55
56
57
58
59
60

- 1
2
3 Tournier J-D, Calamante F, Gadian DG, Connelly A. 2004. Direct estimation of the fiber
4 orientation density function from diffusion-weighted MRI data using spherical
5 deconvolution. *Neuroimage*. 23:1176–1185.
6
7
8
9
10
11 Tyler LK, Chiu S, Zhuang J, Randall B, Devereux BJ, Wright P, Clarke A, Taylor KI. 2013.
12 Objects and Categories: Feature Statistics and Object Processing in the Ventral Stream. *J*
13 *Cogn Neurosci*. 2013 Oct;25(10):1723-35. doi: 10.1162/jocn_a_00419. Epub 2013 May 10.
14
15
16
17
18
19
20 Van Essen DC, Maunsell JHR. 1983. Hierarchical organization and functional streams in the
21 visual cortex. *Trends Neurosci*. 6:370–375.
22
23
24
25
26 Vartanian O. 2012. Dissociable neural systems for analogy and metaphor: implications for the
27 neuroscience of creativity. *Br J Psychol*. 103:302–316.
28
29
30
31
32 Vigneau M, Beaucousin V, Herve PY, Duffau H, Crivello F, Mazoyer B, Tzourio-mazoyer N.
33 2006. Meta-analyzing left hemisphere language areas : Phonology , semantics , and
34 sentence processing. *Neuroimage*. 30:1414–1432.
35
36
37
38
39
40 Visser M, Jefferies E, Lambon Ralph MA. 2010. Semantic processing in the anterior temporal
41 lobes: a meta-analysis of the functional neuroimaging literature. *J Cogn Neurosci*. 22:1083–
42 1094.
43
44
45
46
47
48
49 Volle E, Gilbert SJ, Benoit RG, Burgess PW. 2010. Specialization of the Rostral Prefrontal
50 Cortex for Distinct Analogy Processes. *Cereb Cortex*. 20:2647–2659.
51
52
53
54
55 Wartenburger I, Heekeren HR, Preusse F, Kramer J, van der Meer E. 2009. Cerebral correlates
56 of analogical processing and their modulation by training. *Neuroimage*. 48:291–302.
57
58
59
60

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3
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7
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41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
- Watson CE, Chatterjee A. 2012. A Bilateral Frontoparietal Network Underlies Visuospatial Analogical Reasoning. *Neuroimage*. 59:2831–2838.
- Wei T, Liang X, He Y, Zang Y, Han Z, Caramazza A, Bi Y. 2012. Predicting Conceptual Processing Capacity from Spontaneous Neuronal Activity of the Left Middle Temporal Gyrus. *J Neurosci*. 32:481–489.
- Wendelken C, Chung D, Bunge SA. 2012. Rostrolateral Prefrontal Cortex : Domain-General or Domain-Sensitive ? *1963*:1952–1963.
- Wendelken C, Nakhachenko D, Donohue SE, Carter CS, Bunge SA. 2008. “Brain is to thought as stomach is to ??”: investigating the role of rostralateral prefrontal cortex in relational reasoning. *J Cogn Neurosci*. 20:682–693.
- Wendelken C, Bunge SA, Carter CS. 2008. Maintaining structured information: an investigation into functions of parietal and lateral prefrontal cortices. *Neuropsychologia*. 46: 665-78.
- Wharton CM, Grafman J, Flitman SS, Hansen EK, Brauner J, Marks A, Honda M. 2000. Toward neuroanatomical models of analogy: a positron emission tomography study of analogical mapping. *Cogn Psychol*. 40:173–197.
- Yuan Z, Qin W, Wang D, Jiang T, Zhang Y, Yu C. 2012. The salience network contributes to an individual’s fluid reasoning capacity. *Behav Brain Res*. 229:384–390.

Tables

Table 1. VBM whole-brain analysis showed positive GM correlations with mean accuracy in Analogy, Match, Find and Apply conditions. Whole-brain analysis on GM volume was conducted to investigate significant results at $p < .001$ uncorrected for multiple comparisons with a minimal cluster size of 100 voxels. All significant positive correlations are reported for each condition with their associated brain regions and BA. The MNI coordinates, P (unc.), T values, and cluster size are reported. aITG = Anterior and Inferolateral part of the Inferior Temporal gyrus; NS: non significant.

Condition	Brain region	BA	MNI coordinates			<i>P</i> (unc.)	<i>T</i> value	Cluster size	Label
			x	y	z				
			Analogy	aITG	20				
	aITG	20	-52	-9	-26	<.001	4.43	175	ITEMP
Match	NS								
Find	NS								
Apply	NS								

Table 2. VBM whole-brain analysis showed negative GM correlations with mean accuracy in Analogy, Match, Find and Apply conditions. Whole-brain analysis on GM volume was conducted to investigate significant results at $p < .001$ uncorrected for multiple comparisons with a minimal cluster size of 100 voxels. All significant negative correlations are reported for each condition with their associated brain regions and BA. The MNI coordinates, P (unc.), T values, and cluster size are reported. MFG = Middle Frontal Gyrus; aFG = Anterior part of the Fusiform Gyrus; IFS: Inferior Frontal sulcus IPL= Inferior Parietal lobule; ITG: Inferior Temporal Gyrus; rIPFC = rostrolateral prefrontal cortex. NS: non significant.

Task	Brain region	BA	MNI coordinates			P (unc.)	T value	Cluster size	Label
			x	y	z				
Analogy	rIPFC	10/46	-40	53	1	< .001	4.097	173	IPOL
	(MFG)								
Match	aFG	20	30	-1	-38	< .001	4.07	165	rTEMPmatch
	ITG	20	51	-15	-35	< .001	3.75	105	
	IPL	2	-46	-28	43	< .001	4.13	221	
Find	IFS	45	-39	38	16	< .001	4.27	120	IPFC
Apply	NS	-	-	-	-	-	-	-	-

Table 3. Linear regression analyses between significant VBM clusters and each experimental condition. In each model, age, genre, education, total GM volume were entered as covariates of non-interest.

Dependant variable:	Predictor: AnalogyApply,	Predictors: AnalogyFind,	Predictors MatchApply	Predictors MatchFind
IPOL volume	F(5,48) = 5.890, $p < .001$, accounted for approximately 32% of the variance IPOL volume was predicted by AnalogyApply score (Beta = -.425, $p = .003$) and to a less extent by age (Beta = -.387, $p = .028$)	F(5,48) = 5.40, $p = .001$, accounted for approximately 30% of the variance IPOL volume was predicted by AnalogyFind score (Beta = -.392, $p = .006$) and to a less extent by age (Beta = -.412, $p = .023$)	F(5,48) = 3.278, $p = .013$, accounted for approximately 18% of the variance IPOL volume was not significantly predicted by MatchApply score (Beta = -.066, $p = .613$) nor by age, gender, education or total GM volume	F(5,48) = 3.255, $p = .013$, accounted for approximately 17.5% of the variance IPOL volume was not significantly predicted by MatchFind score (Beta = -.054, $p = .680$) nor by age, gender, education or total GM volume
rTEMP volume	F(5,48) = 2.283, $p = .061$, accounted for approximately 11% of the variance rTEMP volume was predicted by AnalogyApply score only (Beta = .256, $p = .024$)	F(5,48) = 3.748, $p = .006$, accounted for approximately 21% of the variance rTEMP volume was predicted by AnalogyFind score only (Beta = .503, $p = .001$)	F(5,48) = 1.079, $p = .384$, accounted for approximately 1% of the variance rTEMP volume was not predicted by MatchApply score (Beta < .001, $p = .998$) nor by age, gender, education or total GM volume	F(5,48) = 1.009, $p = .378$, accounted for approximately 1% of the variance rTEMP volume was not predicted by MatchFind score (Beta = .031, $p = .831$) nor by age, gender, education or total GM volume
ImidPFC volume	F(5,48) = 9.207, $p < .001$, accounted for approximately 44% of the	F(5,48) = 13.393, $p < .001$, accounted for approximately 54% of the	F(5,48) = 8.159, $p < .001$, accounted for approximately 40% of the variance	F(5,48) = 11.569, $p < .001$, accounted for approximately 50% of the

	variance lmidPFC volume was predicted by age (Beta = - .506, $p = .002$) but not by AnalogyApply score (Beta = - .205, $p = .098$)	variance lmidPFC volume was predicted by AnalogyFind score (Beta = - .416, $p < .001$), by age (Beta = - .564, $p < .001$) and to a less exetent by total GM volume (Beta = .329, $p = .023$)	lmidPFC volume was predicted by age (Beta = -.480, $p = 0.004$) but not by MatchApply score (Beta = - .012, $p = .912$)	variance lmidPFC volume was predicted by AnalogyFind score (Beta = - .307, $p = .004$), by age (Beta = - .527, $p = .001$)
rTEMPmatch volume	F(5,48) = 1.080, $p = .383$, accounted for approximately 1% of the variance rTEMPmatch volume was not predicted by AnalogyApply score (Beta = - .005, $p = .974$)	F(5,48) = 1.167, $p = .339$, accounted for approximately 1.5% of the variance rTEMPmatch volume was not predicted by AnalogyFind score (Beta = .101, $p = .535$) nor by age, gender, education or total GM volume	F(5,48) = 2.218, $p = .068$, accounted for approximately 10% of the variance rTEMPmatch volume was predicted by MatchApply score (Beta = -.306, $p =$.028) and gender (Beta = -.303, $p =$.033)	F(5,48) = 1.981, $p = .098$, accounted for approximately 8.5% of the variance rTEMPmatch volume was predicted by MatchFind score (Beta = -.275, $p =$.050) and gender (Beta = - .305, $p = .034$)

Table 4. SVC analysis: ROI analysis ($p < .05$ corrected based on Family-wise Error) examining the negative correlation between GM volume in *a priori* defined rIPFC regions (defined from fMRI) and accuracy in tasks conditions (in percent of correct responses). Cluster size, T values, fwe-corrected P values, and cluster size are provided. rIPFC = rostrolateral prefrontal cortex.

ROI	MNI			Condition	p (fwe)	T value	Cluster size
	coordinates (x, y, z)						
left rIPFC	-44	50	-4	Mean Analogy	0.003	4.09	189
				Mean Find	ns	-	-
				AnalogyApply	0.013	3.54	160
				AnalogyFind	0.020	3.35	77
				MatchApply	ns	-	-
				MatchFind	ns	-	-
right rIPFC	+44	50	-4	Mean Analogy	ns	-	-
				Mean Find	ns	-	-
				AnalogyApply	ns	-	-
				AnalogyFind	ns	-	-
				MatchApply	ns	-	-
				MatchFind	ns	-	-

Figure captions

Figure 1: A trial for each condition differentiating between Apply and Find conditions in the Analogy and Match tasks. In each condition, participants were asked to choose from the sets of stimuli on the right (target sets) the one that matched the left set of stimuli (source set) according to four distinct conditions. In the AnalogyApply condition (bottom left), participants had to apply a given analogical relationship to the sets to determine the correct target (here, the relationship consisted of a mirror image between the left and right stimuli (letter “g”) of the source set); the correct response is the bottom right target set), while in the AnalogyFind condition (top left), the relationship had to be found by the participant (here, the relationship is an increase in lightness of the stimuli in the source set; the correct response is the top right target set). In the MatchApply condition (bottom right), participants had to apply a matching rule based on a given perceptual feature between the source and correct target set (in the displayed example, the matching feature is colors, and the correct answer is the top right target set), while in the MatchFind condition (top right), they had to find the matching rule, i.e. the perceptual feature shared between the sets (in the displayed example, the left source set and the top right target set share a common number of stimuli).

All of the displayed examples consist of intradimension analogies. However, the analogy task included both intradimension and cross-dimension analogies, as described in Volle et al. 2010 and illustrated in supplementary Figure S1.

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6 **Figure 2.** Mean Accuracy (in % of correct responses) and reaction times (RT in ms) of each
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8 condition. Overall, participants were significantly more accurate and responded faster in the
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10 Match conditions compared to the Analogy conditions. Error bars indicate standard deviations;
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13 ** indicates the significant difference in accuracy and in RTs between Analogy mean and Match
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15 mean conditions.
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21 **Figure 3. Results from the VBM analyses.**

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23 Significant regions associated with variations in GM volume related to performance are
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25 superimposed on an anterolateral view (top left) and anteroinferior view (top right) of a brain
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27 rendering. The VBM whole-brain analyses identified a left rIPFC region (“lPOL”, in red), in
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29 which GM volume negatively correlated with mean performance on Analogy tasks, a left and
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31 right anterolateral temporal region (in green), in which GM volume positively correlated with
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33 mean performance on Analogy tasks, a left caudal prefrontal region (“lmidPFC”, purple), in
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35 which GM volume negatively correlated with mean performance on Find trials, and a right
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37 anteromedial temporal region (“rTEMPmatch”, in blue), in which GM volume negatively
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39 correlated with mean performance on Match tasks.
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45 GM measures were extracted from each individual VBM preprocessed images and averaged
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47 across voxels within these 4 significant clusters evidenced in the whole-brain analyses.
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49 Performance on each experimental condition was entered as a dependent variable and GM
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51 volume in each region as an independent variable in separate multiple regression models, in
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53 which age, gender, education and total GM volume were covaried out.
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3 Plots between performance on each experimental condition and GM measures within these four
4 regions are displayed: the left rIPFC region ('IPOL', in red), the left lateral PFC region
5 ('lmidPFC', in purple), the right anterolateral temporal region ('rTEMP', in green), and an
6 anteromedial temporal region ("rTEMPmatch", in blue). X axes represent the residuals of
7 accuracy in each experimental condition (AnalogyApply, AnalogyFind, MatchApply, Match
8 Find) and Y axes the residuals of the mean GM volume within each region (IPOL, lmidPFC,
9 rTEMP, rTEMPmatch).

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23 **Figure 4. Connectome of the left rIPFC region (IPOL), left and right ATL (ITEMP and**
24 **rTEMP), rTEMPmatch, and lmidPFC region.** Tracts of IPOL (in red), r/ITEMP (in green),
25 rTEMPmatch (in blue), and lmidPFC (in purple) are superimposed on a transparent brain
26 rendering (left side) and on axial slices showing their anatomical connectivity (right side). The
27 upper part of the figures shows the overlap of IPOL and ITEMPT tracts on the arcuate fasciculus.
28 The inferior part of the figure shows that rTEMP and rTEMPmatch tracts poorly overlap on the
29 ILF, and that lmidPFC has mainly intra-frontal connections. ATR: anterior thalamic radiations;
30 FMT: Frontomarginal fasciculus; IFOF: inferior fronto-occipital fasciculus; ILF: inferior
31 longitudinal fasciculus; FIL: Frontal inferior longitudinal fasciculus.

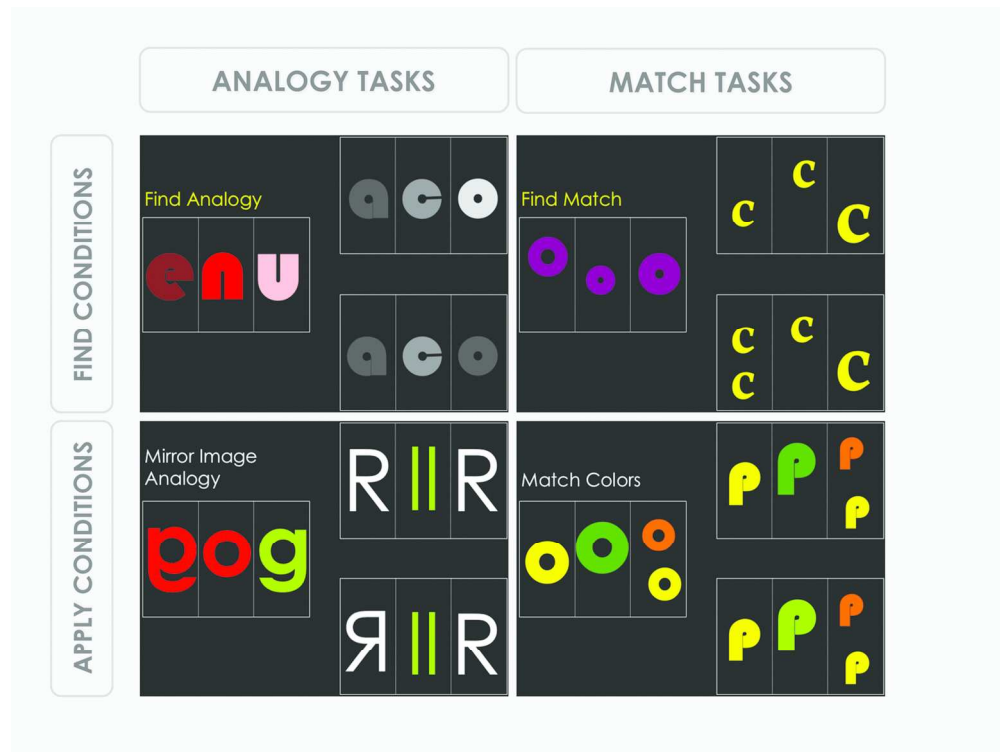


Figure 1. A trial for each condition differentiating between Apply and Find conditions in the Analogy and Match tasks. In each condition, participants were asked to choose from the sets of stimuli on the right (target sets) the one that matched the left set of stimuli (source set) according to four distinct conditions. In the AnalogyApply condition (bottom left), participants had to apply a given analogical relationship to the sets to determine the correct target (here, the relationship consisted of a mirror image between the left and right stimuli (letter "g") of the source set); the correct response is the bottom right target set), while in the AnalogyFind condition (top left), the relationship had to be found by the participant (here, the relationship is an increase in lightness of the stimuli in the source set; the correct response is the top right target set). In the MatchApply condition (bottom right), participants had to apply a matching rule based on a given perceptual feature between the source and correct target set (in the displayed example, the matching feature is colors, and the correct answer is the top right target set), while in the MatchFind condition (top right), they had to find the matching rule, i.e. the perceptual feature shared between the sets (in the displayed example, the left source set and the top right target set share a common number of stimuli). All of the displayed examples consist of intradimension analogies. However, the analogy task included both intradimension and cross-dimension analogies, as described in Volle et al. 2010 and illustrated in supplementary Figure S1.

160x119mm (300 x 300 DPI)

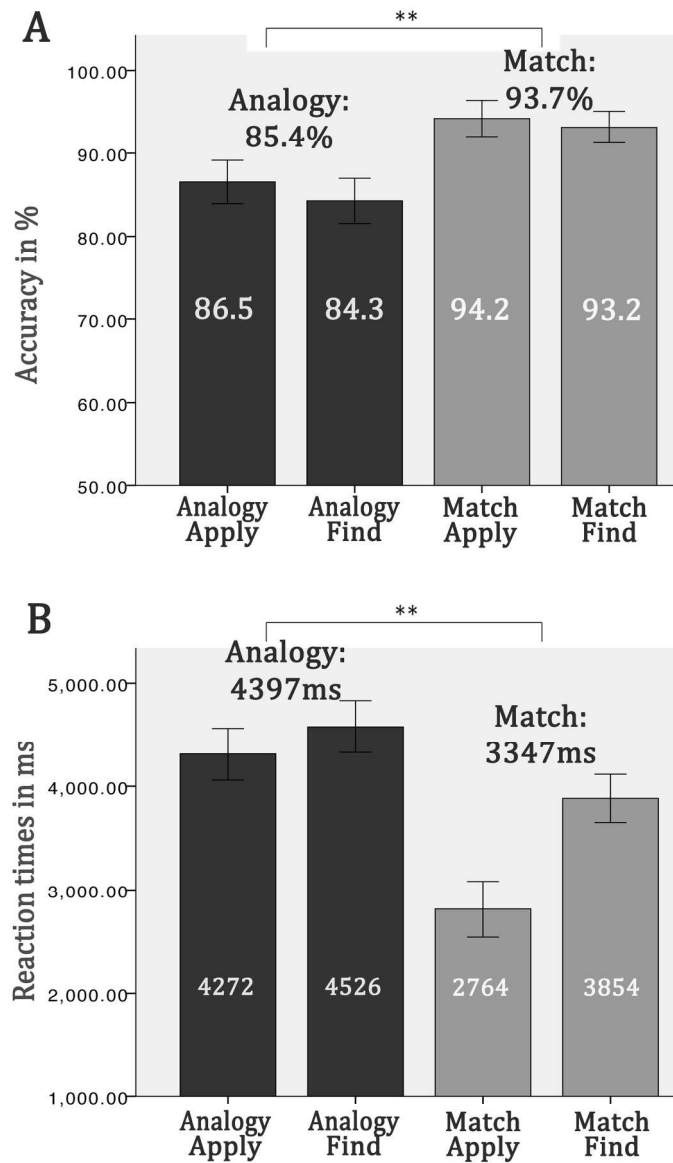


Figure 2. Mean Accuracy (Acc in % of correct responses) and reaction times (RT in ms) of each condition.

Overall, participants were significantly more accurate in the Match conditions compared to the Analogy conditions and responded faster in the Match conditions compared to the Analogy conditions. Error bars indicate standard deviations; ** indicates the significant difference in accuracy and in RTs between Analogy mean and Match mean conditions.

137x235mm (300 x 300 DPI)

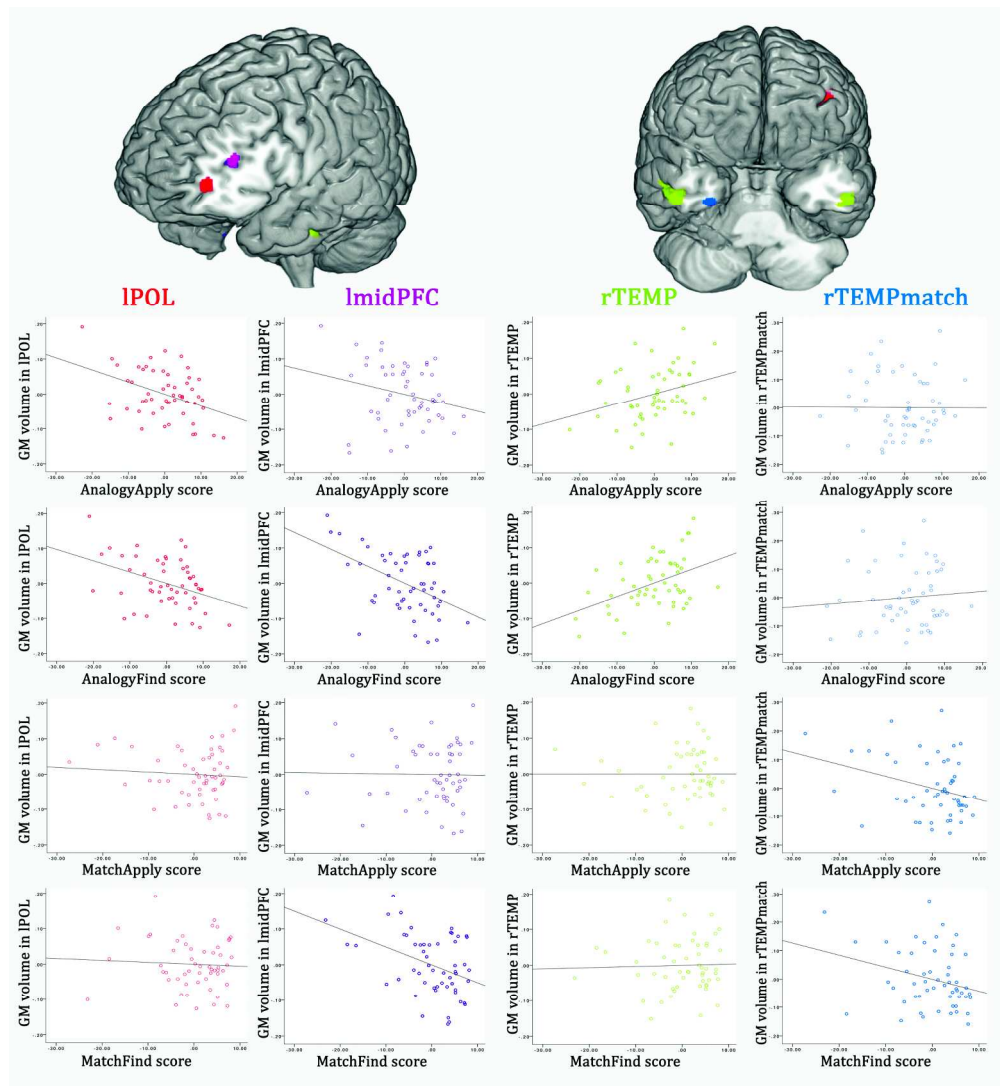


Figure 3. Results from the whole-brain VBM analysis.

Significant regions associated with changes in GM volume related to performance are superimposed on an anterolateral view (left) and anteroinferior view (right) of a brain rendering. The VBM whole-brain analyses identified a left rIPFC region ("IPOL", in red), in which GM volume negatively correlated with mean performance on Analogy tasks, a left and right anterolateral temporal region (in green), in which GM volume positively correlated with mean performance on Analogy tasks, a left caudal prefrontal region ("lmidPFC", purple), in which GM volume negatively correlated with mean performance on Find trials, and a right anteromedial temporal region ("rTEMPmatch", in blue), in which GM volume negatively correlated with mean performance on Match tasks.

GM measures were extracted from each individual VBM preprocessed images and averaged across voxels within the significant clusters evidenced in the whole-brain analysis. Performance on each experimental condition was entered as a dependent variable and GM volume in these four regions as an independent variable in separate multiple regression models, in which age, gender, education and total GM volume were covaried out.

Plots between performance on each experimental condition and GM measures within these four regions are displayed: the left rIPFC region ('IPOL', in red), the left lateral PFC region ('lPFC', in purple), the right anterolateral temporal region ('rTEMP', in green), and an anteromedial temporal region ('rTEMPmatch', in

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blue). X axes represent the residuals of accuracy in each experimental condition (AnalogyApply, AnalogyFind, MatchApply, Match Find) and Y axes the residuals of the mean GM volume within each cluster observed in the whole-brain analysis (IPOL, lmidPFC, rTEMP, rTEMPmatch).

199x216mm (300 x 300 DPI)

For Peer Review

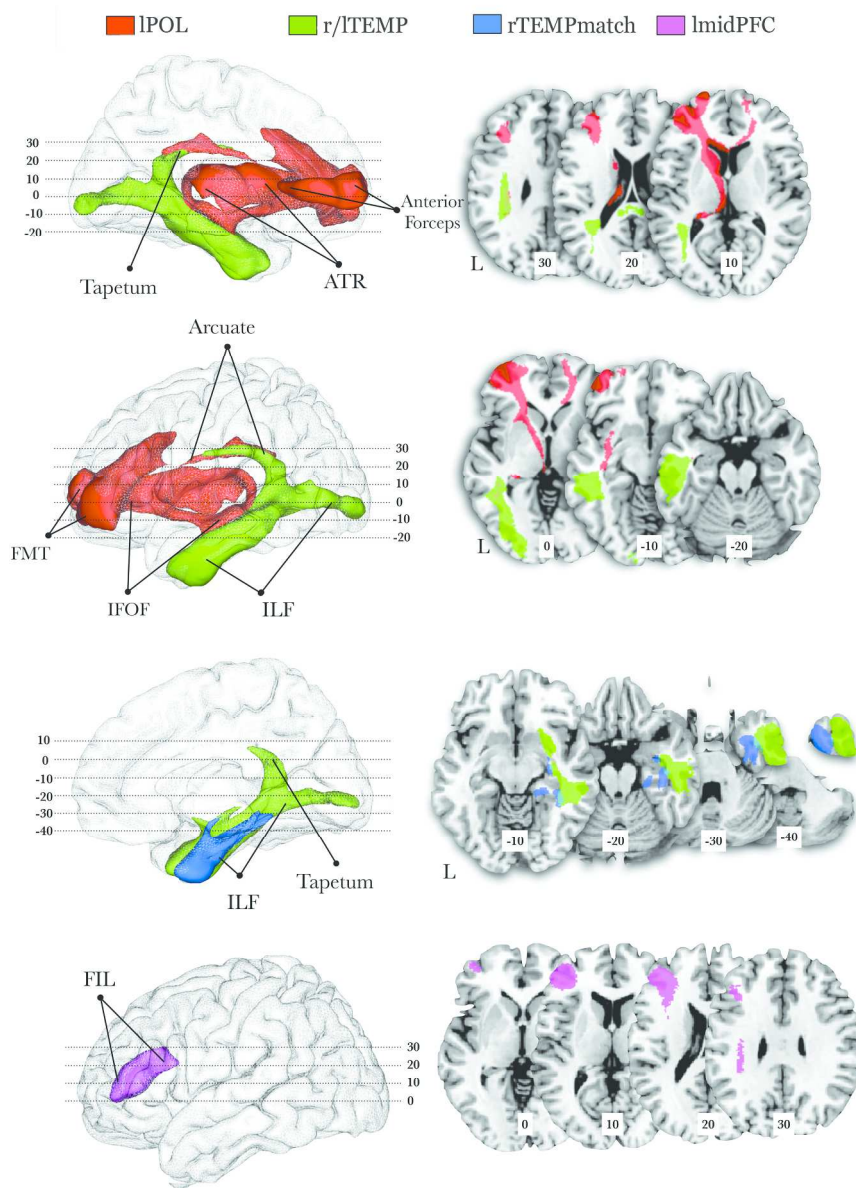


Figure 4. Connectome of the left rIPFC region (IPOL) left and right ATL (ITEMP and rTEMP) and lmidPFC region. Tracts of IPOL (in red), r/ITEMP (in green) and lmidPFC (in purple) are superimposed on a transparent brain rendering (left side) and on axial slices of the anatomical connectivity (right side). The upper part of the figures shows the overlap of IPOL and ITEMPT tracts on the arcuate fasciculus. rTEMP and rTEMPMATCH tracts poorly overlap on the ILF. lmidPFC had mainly intrafrontal connections. ATR: anterior thalamic radiations; FMT: Frontomarginal fasciculus; IFOF: inferior fronto-occipital fasciculus; ILF: inferior longitudinal fasciculus; FIL: Frontal inferior longitudinal fasciculus.

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Supplementary figure S1: Example of intra- and cross-dimension analogy tasks using the same analogy schema “symmetry”, and example of the match task.

Intra-dimension (left column) and cross-dimension (middle column) analogies used the same relational concepts. The figure displays examples of analogy trials using the schema “symmetry”. In the intra-dimension task, the analogy concerned the same dimension in both the source and target sets (e.g., symmetry of the letter identity (top left), of colors (middle left) or the size (bottom left). In the cross-dimension task, the analogy concerned different dimensions (for instance, symmetry of size of the stimuli in the source and symmetry of color in the target stimuli – bottom of the middle column of the figure). The analogy schemas could also concern either the identity of figures, the number of stimuli, their lightness, or their texture. The features of the stimuli that were non relevant for the analogy schema (size, colors, identity, position, texture, or number) varied between source and target in order to avoid perceptual matching. In addition to “symmetry”, there were 5 other different analogy schemas to discover in intra- and cross-dimension analogies, which were not used during the training. These schemas were either visuospatial or mathematical. They could be verbalized as “progressive increase of a feature across the 3 stimuli in the set,” “mirror image,” “the first plus the second gives the third stimulus,” “the first minus the second gives the third stimulus,” “the last is a multiple of the first.”

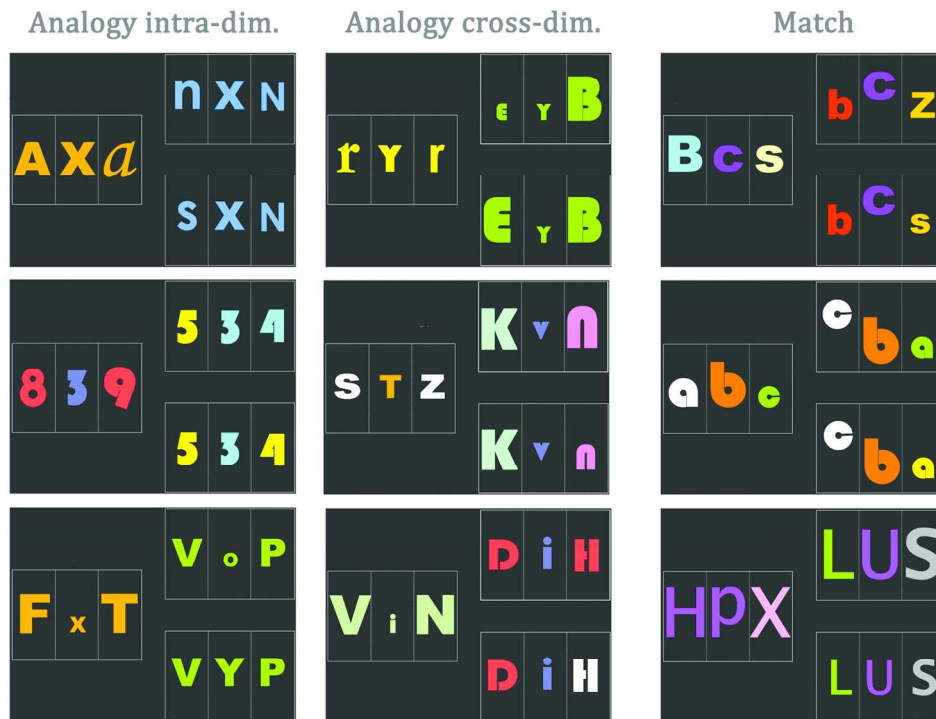
In the Match task, there was one matching feature (i.e. letter identity of the stimuli (top right), identity of colors (middle right), or same size (bottom right) in the displayed trials, while the other features were distractors.

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6 **Supplementary figure S2: Comparison of the whole brain VBM results within rIPFC and**
7 **peak maxima observed in previous fMRI studies.**
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11 Overlap of the current VBM result (whole-brain correlation between Analogy ability and GM
12 volume in the left rIPFC, in red) with regions previously found associated with analogy in
13 functional imaging (in cyan: activation maxima associated with analogy in Volle et al. (2010)
14 fMRI study; in yellow: cluster Maxima observed in Vartanian's metaanalysis (2012) of
15 analogical reasoning).
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29 **Supplementary figure S3: Comparison of the whole brain VBM results within ATL and**
30 **peak maxima observed in previous functional imaging and TMS studies**
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34 Overlap of the current VBM result (whole-brain correlation between Analogy ability and GM
35 volume in bilateral ATL, in green) with regions previously found associated with a semantic
36 memory hub in TMS studies (Lambon Ralph et al 2009, in purple, coordinates -53, 4, -32 and 52,
37 2, -28). L: left side.
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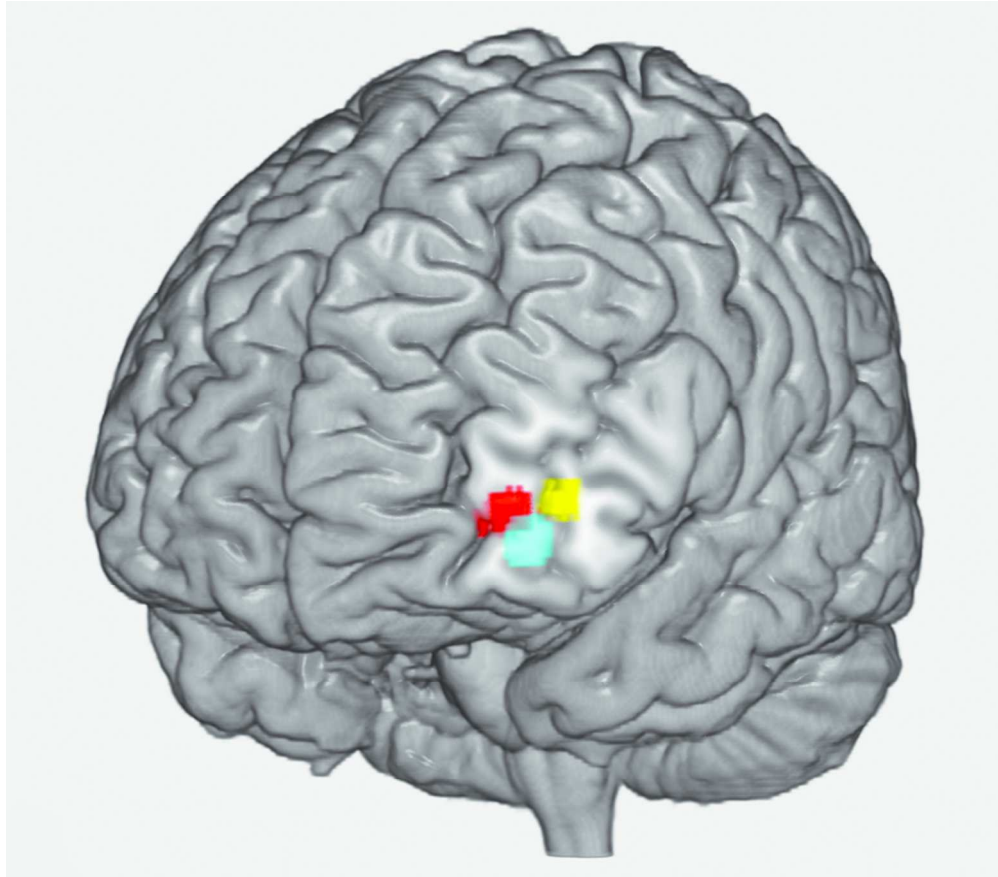


Example of intra- and cross-dimension analogy task using the same analogy schema "symmetry" and of match task.

Intradimension (left column) and cross-dimension (middle column) analogies used the same relational concepts. The figure displays examples of analogy trials using the schema "symmetry". In the intradimension task, the analogy concerned the same dimension in both the source and target sets (e.g., symmetry of the letter identity (top left), of colors (middle left) or the size (bottom left)). In the cross-dimension task, the analogy concerned different dimensions (for instance, symmetry of size of the stimuli in the source and symmetry of color in the target stimuli – bottom of the middle column of the figure). The analogy schemas could also concern either the identity of figures, the number of stimuli, their lightness, or their texture. The features of the stimuli that were non relevant for the analogy schema (size, colors, identity, position, texture, or number) varied between source and target in order to avoid perceptual matching. In addition to "symmetry", there were 5 other different analogy schemas to discover in intra- and cross-dimension analogies, which were not used during the training. These schemas were either visuospatial or mathematical. They could be verbalized as "progressive increase of a feature across the 3 stimuli in the set," "mirror image," "the first plus the second gives the third stimulus," "the first minus the second gives the third stimulus," "the last is a multiple of the first."

In the Match task, there was one matching feature (i.e. letter identity of the stimuli (top right), identity of colors (middle right), or same size (bottom right) in the displayed trials, while the other features were distractors.

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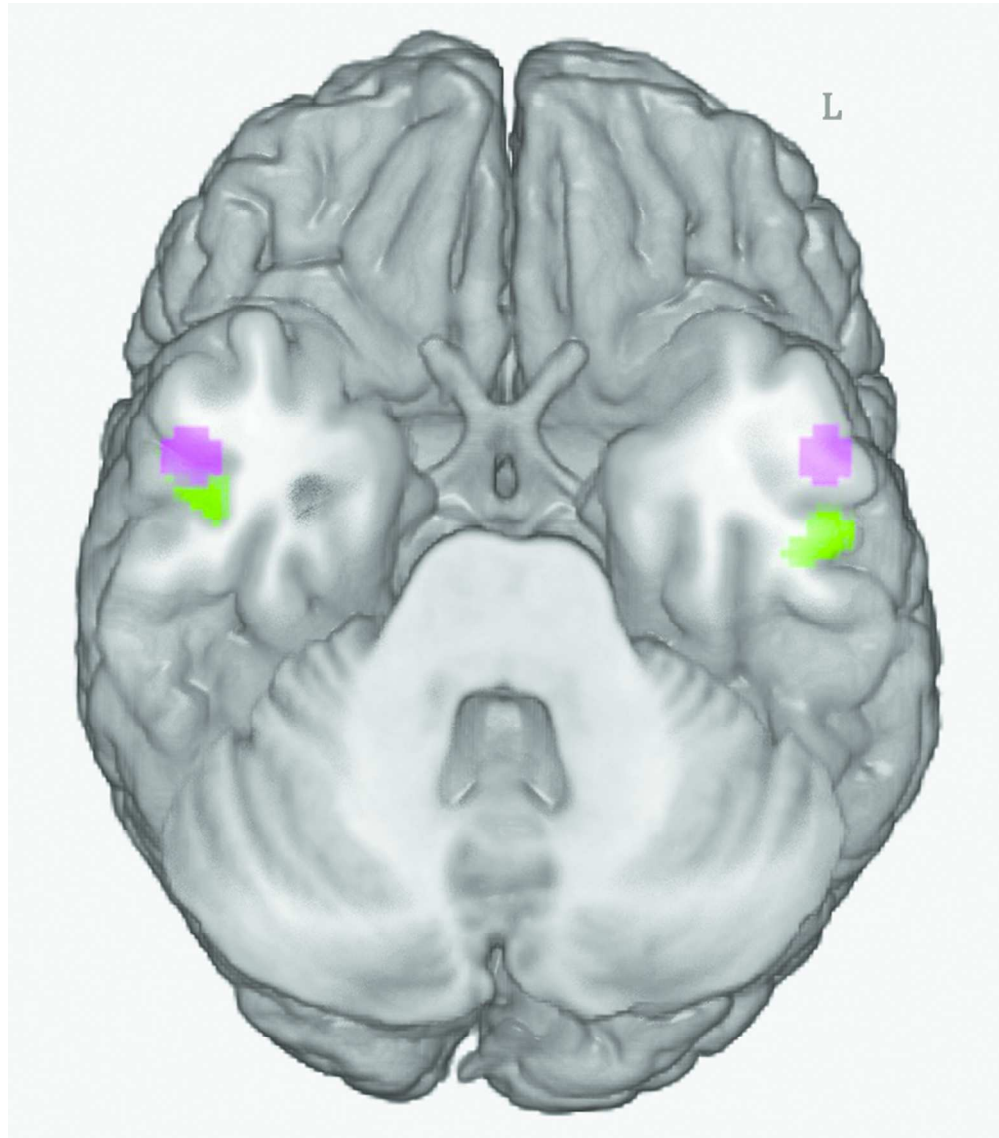


Comparison of the whole brain VBM results within rIPFC and peak maxima observed in previous results fMRI studies

Overlap of the current VBM whole brain correlation with Analogy ability and GM in the left rIPFC with previous regions found associated with analogy in functional imaging.

red: current VBM region showing a negative correlation to analogy in the whole-brain analysis; cyan: Activation maxima associated with analogy in Volle et al 2010 fMRI study; yellow: Cluster Maxima observed in Vartanian's metaanalysis (2012) of analogical reasoning.

80x70mm (300 x 300 DPI)



Comparison of the whole brain VBM results within ATL and peak maxima observed in previous results functional imaging and TMS studies
Overlap of the current VBM whole-brain correlation with Analogy ability and GM in the anterior temporal region with previous regions found associated with a semantic memory hub in TMS studies (Lambon Ralph et al 2009). Green: current VBM region showing a positive correlation with analogy in the whole-brain analysis; purple: Lambon Ralph 2009 sites of TMS stimulation evoking semantic deficits (coordinates -53, 4, -32 and 52, 2, -28). L: left side.

80x90mm (300 x 300 DPI)