

Reasoning, Models, and Images: Behavioral Measures and Cortical Activity

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Abstract

■ The goal of this study was to investigate the neurocognitive processes of mental imagery in deductive reasoning. Behavioral studies yielded four sorts of verbal relations: (1) visuospatial relations that are easy to envisage both visually and spatially; (2) visual relations that are easy to envisage visually but hard to envisage spatially; (3) spatial relations that are hard to envisage visually but easy to envisage spatially; and (4) control relations that are hard to envisage both visually and spatially. In three experiments, visual relations slowed the process of reasoning in comparison with control relations, whereas visuospatial and spatial relations yielded inferences comparable to those of control relations. An experiment using functional magnetic resonance imaging showed that in the

absence of any correlated visual input (problems were presented acoustically via headphones), all types of reasoning problems evoked activity in the left middle temporal gyrus, in the right superior parietal cortex, and bilaterally in the precuneus. In the prefrontal cortex, increased activity was found in the middle and inferior frontal gyri. However, only the problems based on visual relations also activated areas of the visual association cortex corresponding to V2. The results indicate that cortical activity during reasoning depends on the nature of verbal relations. All relations elicit mental models that underlie reasoning, but visual relations in addition elicit visual images. This account resolves inconsistencies in the previous literature. ■

... *it is clear that imagery plays a key role in reasoning.*
(Kosslyn, 1994, p. 404)

INTRODUCTION

Mental imagery has been studied extensively from the standpoint of memory (Block, 1981; Kosslyn, 1980, 1994), but its role in human reasoning remains unclear. If you ask people how they reason, many of them say that they rely on visual images. For instance, with a problem such as

Ann is taller than Bert.
Bert is taller than Cath.
Does it follow that Ann is taller than Cath?

they say that they formed a mental picture in their “mind’s eye” and then looked at this picture to see that Ann is taller than Cath. Reasoning does seem inextricably linked with seeing in the “mind’s eye.” However, does this experience of mental imagery reflect the real nature of underlying mental representations? Some theorists such as Kosslyn say “yes” (see the epigraph to this article), whereas other theorists such as Pylyshyn say “no” (see Pylyshyn, 1981, in press). The aim of this

article is to make progress towards resolving the controversy—at least as it applies to reasoning—and at the same time to examine the neuronal processes that underlie reasoning. The article begins with a description of inferences based on relations and a summary of the apparently inconsistent effects of the imageability of relations. Next, it describes some behavioral studies with such inferences in which we manipulated the nature of the relations. The article reports an functional magnetic resonance imaging (fMRI) study of the same problems that elucidates the relation among images, models, and cortical activity. Finally, it shows how the findings resolve the inconsistencies in the literature, and it draws general conclusions about the role of mental representations in deductive reasoning.

Our research concerns an essential sort of reasoning: deductive reasoning (Rips, 1994). A valid deduction, by definition, yields a conclusion that must be true granted that its premises are true. In contrast, inductive reasoning is not valid, and so an induction from true premises may or may not yield a true conclusion. Consider, for example, the preceding problem. To a logician, the inference yields a transitive conclusion, which depends for its validity on a missing premise: For any x , y , and z , if x is taller than y and y is taller than z , then x is taller than z . Psychologists have studied such inferences, which they refer to as “linear syllogisms” or “three-term series” problems, and they have proposed various

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accounts of how individuals solve them (see, e.g., Evans, Newstead, & Byrne, 1993).

A pioneering study of three-term series inferences was carried out by De Soto, London, and Handel (1965). They argued that reasoners represent the entities in the premise in a visual image and then “read off” the answer by inspecting this image. On this account, problems of the form

A is shorter than B.
B is shorter than C.

are harder than those in which the relation is “taller than,” because reasoners have to build up a representation in an order that is unnatural, working from the shortest individual up to the tallest individual. Huttenlocher (1968) similarly argued that reasoners imagine a spatial array of objects, and that it is easier to move an entity into the array if it is referred to by the subject of the sentence. In contrast, Clark (1969) proposed a linguistic account of the difficulty of relational inferences. Linguistically speaking, a relation such as “taller than” is lexically unmarked: It tells you nothing, for example, about the absolute height of the two entities. But, a relation such as “shorter than” is lexically marked: It tells you that both entities are short by the standards of the class to which they belong. Marked terms, Clark argued, take longer to process, and so they impede the process of reasoning. The controversy between these two accounts was never properly resolved, perhaps because both of them could be right (see Johnson-Laird, 1972). Marked terms could take longer to understand than unmarked terms, but, once understood, they could be used to construct spatial representations of the problems.

If reasoning depends on constructing a visual image, then it should be easier to reason with materials that can be readily visualized. However, if reasoning depends on some other medium of representation, such as sentences in a mental language, then imageability might have no effect on the process. So, does imageability improve reasoning? The literature fails to yield a clear-cut answer. On the one hand, Shaver, Pierson, and Lang (1975) reported that relational reasoning depends on the ease of visualizing the materials, instructions to form images, and the participants’ ability to form images. In addition, Clement and Falmagne (1986) found that materials rated as easy to imagine led to fewer errors in reasoning. On the other hand, several studies have failed to detect any effect of imageability on reasoning. Sternberg (1980) found no difference in accuracy between problems that were easy to visualize and problems that were hard to visualize, and no reliable correlation between reasoning and imaging abilities. Richardson (1987) reported that reasoning with visually concrete problems was no better than reasoning with abstract problems. Similar results have been reported by Johnson-Laird, Byrne, and Tabossi (1989) and Newstead, Pollard, and Griggs (1986).

The inconsistency might arise, as Knauff and Johnson-Laird (2002) have argued, because researchers have overlooked the distinction between visual images and spatial representations. This distinction has been drawn by researchers in cognitive neuroscience (Mellet et al., 2000; Kosslyn, 1994; Rueckl, Cave, & Kosslyn, 1989; Ungerleider & Mishkin, 1982), psycholinguists and linguists (e.g., Landau & Jackendoff, 1993), and cognitive psychologists (e.g., Johnson-Laird, 1998). It is supported by investigations of brain-damaged patients (e.g., Newcombe, Ratcliff, & Damasio, 1987), by functional brain imaging (e.g., Smith et al., 1995), and by behavioral experiments on working memory (see Logie, 1995). Visual images are structurally similar to real visual perceptions, and they can represent objects, their colors and shapes, and the metrical distances between them. They have a limited resolution, but they can be scanned and mentally manipulated in other ways (see, e.g., Finke, 1989; Kosslyn, 1980). They can be so similar to real perceptions that they can be confused for them (Johnson & Raye, 1981). In contrast, spatial representations are models of the spatial relations among entities, that is, they represent what things are where. They might also be used to represent abstract relations in a spatial way, e.g., class inclusion can be represented by the spatial inclusion of one area within another. In inferential tasks, spatial models are likely to exclude visual detail, to represent only the information relevant to inference (Johnson-Laird, 1998), and to take the form of multidimensional arrays that maintain ordinal and topological properties (Knauff, 1999; Knauff, Rauh, Schlieder, & Strube, 1998). In sum, visual images represent information in a modality-specific format, whereas spatial models are abstract and not restricted to a specific modality. We can therefore distinguish between two hypotheses about reasoning. According to the imagery hypothesis, reasoning depends on visual images, and so brain regions that mediate them should be active during reasoning. According to the spatial hypothesis, reasoning depends on spatial models, and so brain regions that mediate them should be active during reasoning, but specific materials that evoke imagery should in addition activate regions mediating images.

The distinction between visual images and spatial representations was corroborated in norming studies that we carried out to determine whether the ease of visualizing a relation might be independent of the ease of forming a spatial representation of it (Knauff & Johnson-Laird, 2000). The data from these studies yielded four sorts of relations:

1. visuospatial relations that are easy to envisage visually and spatially, such as “above” and “below”
2. visual relations that are easy to envisage visually but hard to envisage spatially, such as “cleaner” and “dirtier”

3. spatial relations that are difficult to envisage visually but easy to envisage spatially, such as “further north” and “further south”

4. control relations that are hard to envisage both visually and spatially, such as “better” and “worse”

These four sorts of relations were used to construct four corresponding classes of inferential problems that were used in the following experiments (the details are described in Methods). All inferences in the experiments used the same nouns (“dog,” “cat,” and “ape”) to minimize differences as a result of anything other than the relations. Here is an example of an inference with a valid conclusion:

The dog is cleaner than the cat.

The ape is dirtier than the cat.

Does it follow:

The dog is cleaner than the ape?

The participants had to decide whether the conclusion followed from the premises. The inferences were matched in all respects except for the nature of the relation. In the behavioral experiments, the problems were presented visually on a computer screen. In the fMRI experiment (and its prior training study), they were presented acoustically via pneumatic headphones to avoid any visual input. Although the inferences based on control relations were treated as a baseline for assessing differences in the neural processing of the four types of problems, brain images recorded during rest intervals were also used to assess the activation evoked by the entire set of reasoning problems. On the neuroanatomical level, the spatial hypothesis predicts that parietal cortical areas underlying spatial working memory, perception, and movement control, should be evoked by reasoning, whereas the imagery hypothesis postulates that areas of the occipital cortex should be evoked by reasoning (e.g., Knauff, Mulack, Kassubek, Salih, & Greenlee, 2002). Hence, the analysis of the imaging data was carried out in two steps. The first step

identified the cortical areas active for reasoning in general (visual, visuospatial, spatial, and control inferences vs. the rest interval). The second step examined differences among the four sorts of relation.

RESULTS

Behavioral

Behavioral Experiments

Table 1 presents the mean latencies for the correct responses and the percentages of correct responses for the four sorts of relational inferences in the three experiments. There was no reliable difference in accuracy depending on the four kinds of relations in any of the experiments (about 80% correct overall). The first two experiments used only visuospatial, visual, and control inferences. The visuospatial inferences tended to be faster than the control inferences, but the difference was not significant. However, the visual inferences were reliably slower than the control inferences (Experiment 1: Wilcoxon test, $z = 3.07$, $p < .0025$; Experiment 2: Wilcoxon test, $z = 2.46$, $p < .025$). The third experiment used all four sorts of inferences, and there was a reliable trend of increasing response latencies of the following order: spatial inferences < visuospatial inferences < control inferences < visual inferences (Page’s $L = 648$, $z = 3.40$, $p < .01$). Again, however, the principal effect was that visual relations slowed reasoning relative to the other three sorts of relation (Wilcoxon test, $z = 2.49$, $p < .01$), which did not differ reliably from each other.

Training Experiment

The participants in the brain scanning study carried out a prior training experiment in which the computer presented the problems acoustically using the same timings as in the scanning study. The results, which are also shown in Table 1, corroborated the previous findings. The inferences were easy (89% correct overall),

Table 1. Mean Response Latencies (in sec) and the Percentages of Correct Responses (in parentheses) to the Four Sorts of Inference in Three Behavioral Experiments (from Knauff & Johnson-Laird, 2002), the Training Experiment, and the Scanning Experiment

	<i>Visual Inferences</i>	<i>Control Inferences</i>	<i>Visuospatial Inferences</i>	<i>Spatial Inferences</i>
Experiment 1	2.65* (86)	2.38 (92)	2.20 (90)	–
Experiment 2	3.37* (77)	2.64 (67)	2.46 (68)	–
Experiment 3	4.48* (73)	3.81 (78)	3.74 (72)	3.52 (72)
Training experiment	2.6 (93)	2.3 (91)	2.3 (88)	2.2 (83)
Scanning experiment	2.1 (71)	2.0 (95)	2.0 (77)	2.0 (94)

Experiments 1 and 2 used only three sorts of inferences.

*Statistically significant difference from the control inferences.

Table 2. Activation Clusters in Contrasts between the Images Acquired during the Four Sorts of Inferences and Those Acquired during the Rest Intervals

Location	z Score ^a	Size (Voxels)	Talairach Coordinates ^a		
			x	y	z
<i>Visuospatial versus Rest Interval</i>					
Parietal					
Left superior parietal gyrus (BA 7)	4.80	59***	-21	-61	56
Right superior parietal gyrus (BA 7)	5.36	24***	27	-56	55
Temporal					
Left middle temporal gyrus (BA 21)	6.96	64***	-62	-32	2
Frontal					
Left middle frontal gyrus (BA 11)	4.81	35***	-45	34	-14
<i>Visual versus rest interval</i>					
Parietal					
Right superior parietal gyrus (BA 7)	4.95	14 ^b	27	-56	55
Temporal					
Left middle temporal gyrus (BA 21)	5.65	80***	-59	-12	-7
Right middle temporal gyrus (BA 21)	4.25	17*	59	-6	-5
Frontal					
Left middle frontal gyrus (BA 11)	5.03	29**	-42	34	-14
Left middle frontal gyrus (BA 9)	4.95	22**	-50	25	26
Right middle frontal gyrus (BA 6)	4.17	15*	30	-3	50
<i>Spatial versus rest interval</i>					
Parietal					
Left precuneus (BA 7)	4.23	17*	-21	-58	55
Right precuneus (BA 7)	5.29	23**	24	-56	53
Temporal					
Left middle temporal gyrus (BA 21)	4.79	18***	-59	-29	1
Frontal					
Left inferior frontal gyrus (BA 47)	4.70	18*	-48	40	-15
<i>Control versus rest interval</i>					
Parietal					
Left precuneus (BA 7)	5.51	58***	-18	-65	47
Right superior parietal gyrus (BA 7)	4.29	34**	27	-52	63
Temporal					
Left middle temporal gyrus (BA 21)	4.58	67***	-59	-9	-5

^aLocation, z score, and Talairach coordinates refer to the peak voxel of the cluster. The correspondence of this voxel to BA is established only when applicable. This localization can be made only at the level of the whole cluster.

^bThis cluster was added to the table, although it is one voxel below the threshold for significance at the cluster level ($p = .056$). Note that precisely this area is active in the conjunction analysis presented in Table 3.

* $p < .05$, corrected for multiple comparisons across the whole brain.

** $p < .01$, corrected for multiple comparisons across the whole brain.

*** $p < .001$, corrected for multiple comparisons across the whole brain.

and there was no significant difference in accuracy across the four sorts of problems. There was also no significant difference between accepting valid conclusions (87% correct) and rejecting invalid conclusions (90% correct), and so we pooled the results from the two conditions. The visual inferences were slower than the control inferences, whereas the visuospatial and spatial inferences did not differ from the control problems. However, owing to the small number of participants, none of the differences were significant (Friedman nonparametric analysis of variance, $F = .23$, $p = .97$; this test assesses differences in a within-subjects design, see Siegel & Castellan, 1988).

Imaging Experiment

Table 1 presents the percentages of correct responses and their mean latencies in the imaging experiment. The participants' accuracy was in the same range (84% correct overall) as in the previous behavioral experiments. However, the response latencies showed the same pattern as before: Correct responses had a tendency to be slower for the visual problems than for the other sorts of problems, though the difference was not statistically significant (Friedman analysis of variance, $F = 4.64$, $p = .20$).

Brain Imaging

Table 2 summarizes the results of the comparisons of brain activation during the four sorts of inference and during the rest intervals. In comparison with the rest intervals, each of the four sorts of inference activated the left middle temporal gyrus corresponding to Brodmann's area (BA) 21 and portions of the right parietal cortex (superior gyrus or precuneus; BA 7). The visuospatial, spatial, and control inferences also activated similar areas: left superior parietal gyrus or precuneus (BA 7). The visuospatial and visual inferences additionally activated areas of the middle frontal gyrus (BA 11), while the spatial inferences also activated the left inferior frontal gyrus (BA 47). Apart from BA 11, however, the visual inferences distinctly activated quite different regions: left and right middle frontal gyrus (BA 9 and 6).

In the light of these patterns of activation, we calculated an additional contrast that directly examined the overlap in the activation patterns from the four sorts of inferences. This contrast displays the areas active during all four sorts of inferences, but it does not display any areas that yield a significant activation for only one of the four types of inference (see Methods for a description of the procedure). Table 3 and Figure 1 present the active areas in this contrast. As the single contrasts already suggested, all four sorts of inference led to activity in the

Table 3. Activation Clusters in a Contrast between the Images Acquired during All Four Sorts of Inference and Those Acquired during the Rest Intervals

Location	z Score	Size (Voxels)	Talairach Coordinates		
			x	y	z
<i>All inferences versus rest interval</i>					
Parietal					
Left precuneus (BA 7)	6.00	25**	-18	-58	55
Right precuneus (BA 7)	4.72	16*	15	-65	45
Right superior parietal gyrus (BA 7)	6.59	41***	27	-56	55
Temporal					
Left middle temporal gyrus (BA 21)	7.40	46***	-59	-32	2
Right middle temporal gyrus (BA 21)	5.16	25**	65	-15	-2
Left superior temporal gyrus (BA 38)	4.57	15*	-54	14	-21
Frontal					
Left middle frontal gyrus (BA 46)	5.59	19*	-51	25	24
Left inferior frontal gyrus (BA 47)	5.16	27**	-48	43	-15
Right middle frontal gyrus (BA 6)	4.69	15*	30	-1	47

The clusters were masked inclusively by the comparisons: visuospatial-versus-rest, visual-versus-rest, spatial-versus-rest, and control-versus-rest intervals.

* $p < .05$, corrected for multiple comparisons across the whole brain.

** $p < .01$, corrected for multiple comparisons across the whole brain.

*** $p < .001$, corrected for multiple comparisons across the whole brain.

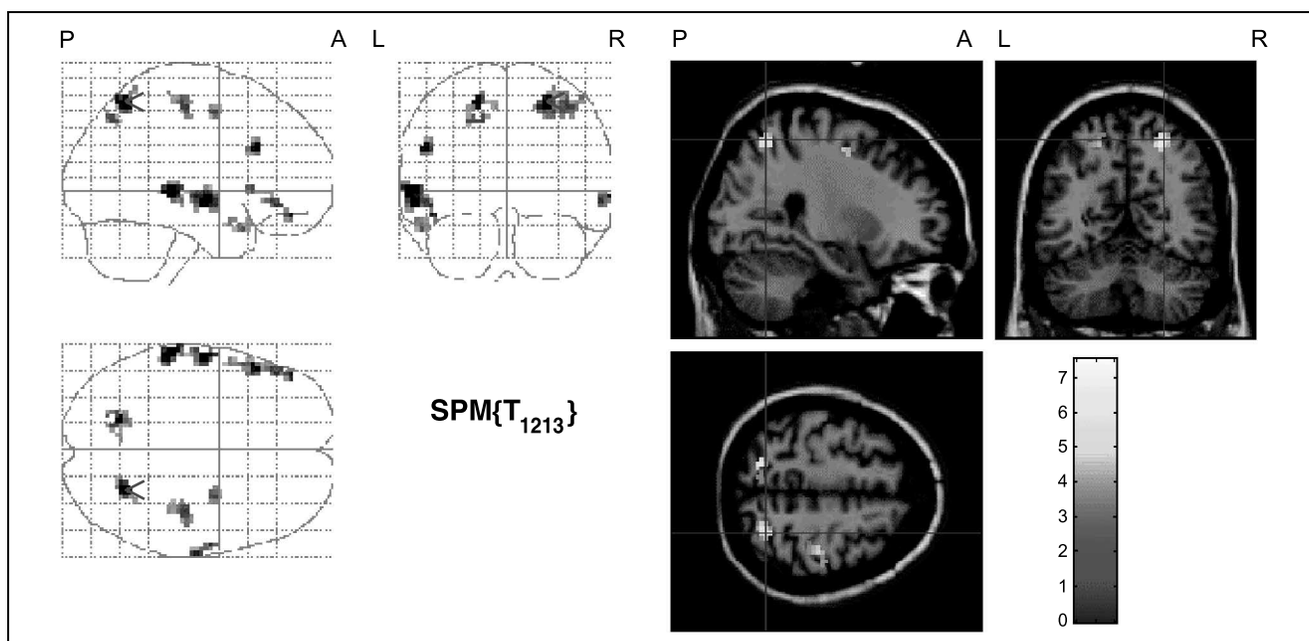


Figure 1. Reasoning with all four sorts of relation activated bilateral parietal cortex. The figure shows the SPM(Z)s of the activation common to contrasts between the four relations and the rest interval. In the left-hand pictures, the z values were transferred to an arbitrary gray scale, and projected onto sagittal, coronal, and transverse sections of a standard brain template. In the right-hand pictures, the z values were plotted according to the color scale given right of each transverse section (L = left hemisphere; R = right hemisphere; A = anterior end; P = posterior end). Crosshairs are positioned in the local peak voxel for the respective contrast and brain area (see Table 3 for other activations). All clusters are significant at $p < .05$ (cluster-level inference) corrected for multiple comparisons across the whole brain.

bilateral precuneus (BA 7), right superior parietal gyrus (BA 7), bilateral middle temporal gyrus (BA 21), left superior temporal gyrus (BA 38), left inferior and middle frontal gyri (BA 47, 46) corresponding to dorsolateral prefrontal cortex (DLPFC), and middle frontal gyrus (BA 6) corresponding to premotor cortex. Owing to the increased statistical power of this contrast, these clusters were larger and had higher z values than those in the single contrasts.

The main aim of the brain imaging study was to determine whether the four sorts of reasoning problems activated different regions of the brain. The control inferences were identical to the other inferences in all respects except that the relations in them are neither easy to visualize nor easy to envisage spatially. Hence, we used them as a baseline in separate contrasts with each of the three other types of inference: visuospatial, spatial, and visual. Table 4 summarizes the results. There were no reliable differences in activation between the control inferences and the visuospatial and spatial inferences. However, in comparison to the control inferences, the visual inferences led to additional activation in an area that covers parts of the visual association cortex corresponding to V2 (BA 18) and the precuneus (BA 31). To determine the location of the activation more precisely, we counted the exact number of voxels falling into each of these areas: 11 of the 33 active voxels were in BA 18, 17 were in BA 31, and the remainder were in BA 17. Surprisingly, this contrast also revealed increased activation in the insula (BA 13).

DISCUSSION

We used behavioral and imaging experiments to examine the cognitive and neural processes underlying reasoning based on various sorts of relations. In what

Table 4. Activation Clusters in Contrasts between the Control Inferences and the Three Other Sorts of Inference

Location	z Score	Size (Voxels)	Talairach Coordinates		
			x	y	z
<i>Visual versus control inferences</i>					
Occipito-parietal					
Right precuneus (BA 18/31)	4.45	33**	12	-72	26
Sublobar					
Left insula (BA 13)	4.68	17*	-30	-31	21
<i>Spatial versus control inferences</i>					
No additional activity measured					
<i>Visuospatial versus control inferences</i>					
No additional activity measured					

* $p < .05$, corrected for multiple comparisons across the whole brain.

** $p < .01$, corrected for multiple comparisons across the whole brain.

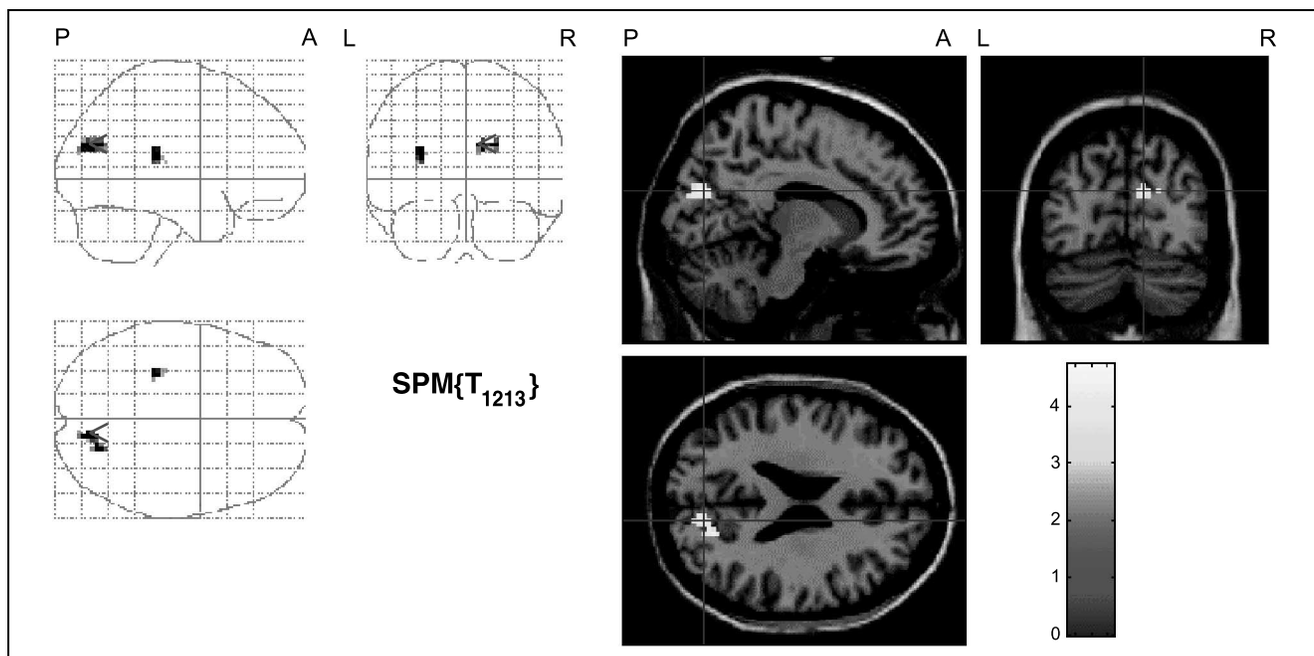


Figure 2. Reasoning about visual relations, that is, those that are easy to visualize but hard to envisage spatially, additionally activated areas in the secondary visual cortex corresponding to V2. The figure shows the SPM(Z)s of the activation from the contrast between visual problems and control problems. In the left-hand pictures, all z values were transferred to an arbitrary gray scale, and projected onto sagittal, coronal, and transverse sections of a standard brain template. In the right-hand pictures, z values are plotted according to the color scale given right of each transverse section (L = left hemisphere; R = right hemisphere; A = anterior end; P = posterior end). Crosshairs are positioned in the local peak voxel for the respective contrast and brain area. All clusters are significant at $p < .05$ (cluster-level inference) corrected for multiple comparisons across the whole brain.

follows, we sketch how the behavioral findings resolve the inconsistency in the previous literature about whether imageability enhances reasoning. We then use the brain imaging data to elucidate the nature of mental representations. Finally, we draw some conclusions about mental imagery, deductive reasoning, and working memory.

The experiments yielded a striking result contrary to the orthodox hypothesis that visual images facilitate reasoning. In fact, they impeded reasoning. They slowed it down in comparison with the other sorts of relations: visuospatial, spatial, and control relations, which tend not to differ reliably in the time that they require for reasoning. The most plausible explanation for this pattern of results is that reasoning is normally based on spatial or abstract mental models. Visual relations, however, spontaneously elicit imagery that is not pertinent to reasoning, and it accordingly takes longer to recover a representation appropriate for reasoning. It is conceivable that all reasoning is based on spatial models, which are pressed into service even for nonspatial relations. It follows from this hypothesis that spatial relations should be dealt with fastest of all. Experiment 3 and the training experiment both showed a trend in this direction (see Table 1), but it is hardly robust.

Previous studies of imagery and reasoning have led to inconsistent results. But, as we argued in the Introduction, these studies have overlooked the distinction between visual and spatial representations. Indeed, the

studies that produced a facilitating effect of “imagery” tended to use materials that were easy to represent spatially, whereas the studies that found no such effect tended to use materials that evoked visual imagery (for details, see Knauff & Johnson-Laird, 2002).

Our brain imaging data provide more evidence for a distinction between visual and spatial representations in reasoning. Our first discovery was that reasoning in general evoked activity in the superior parietal cortex and the precuneus. The parietal cortex is supposed to play a major role in spatial processing and in the integration of sensory information from all modalities into egocentric spatial representations (Xing & Andersen, 2000; Bushara et al., 1999; Andersen, 1997; Andersen, Snyder, Bradley, & Xing, 1997; Colby & Duhamel, 1996; Kolb & Wishaw, 1996). Activation in this area is also interpreted as indicating the use of spatial working memory (Oliveri et al., 2001; Postle, Berger, & D’Esposito, 1999; Smith & Jonides, 1998; Baker, Frith, Frackowiak, & Dolan, 1996). A recent model of the functional network underlying spatial cognition, primarily in navigation, treats parieto-occipital regions as implicated in computing head-centered representations to produce spatial representations of the environment that are held in the precuneus (Burgess, Maguire, Spiers, & O’Keefe, 2001; Maguire, 2001). Previous brain imaging studies of reasoning have similarly implied that parietal cortex plays a key role in reasoning based on mental models, because of their abstract spatial nature (see Knauff et al., 2002; Goel & Dolan, 2001; but cf. Goel, Gold, Kapur, & Houle, 1998).

Our second discovery was that only reasoning based on visual relations evoked additional activity in visual association cortex, V2 (see also the results from a previous fMRI study of visual imagery conducted by our group: Knauff, Kassubek, Mulack, Greenlee, 2000). This region of occipital cortex seems likely to play a key role in visual imagery. It is consistently activated when participants have to visualize objects based on verbal descriptions or previous experience (e.g., Kosslyn et al., 1999; D'Esposito et al., 1997; Kosslyn, Thompson, & Alpert, 1997; Kosslyn, Thompson, Kim, & Alpert, 1995; Mellet, Tzourio, Denis, & Mazoyer, 1995; Mellet et al., 1996; Roland & Gulyás, 1995). Lesions in this region have a devastating effect on the ability to form visual mental images (e.g., Ishai, Ungerleider, Martin, & Haxby, 2000; Farah, 1995).

The contrast between problems based on visual relations and the rest intervals failed to show activation of secondary visual cortex. This failure appears to weaken our interpretation of the results. However, the lack of such activation may merely reflect the inappropriate use of rest intervals as a subtractive baseline in brain imaging studies. Several recent studies have underlined this point, because they have shown that rest intervals yield considerable neuronal activity in regions underlying working memory (Gusnard, Raichle, & Raichle, 2001; Mazoyer et al., 2001; Newman, Twieg, & Carpenter, 2001). This activity may be a default mode of brain functioning or a peaking of task-irrelevant thoughts in the breaks between experimental problems. Hence, the lack of activity in visual cortex in this contrast may merely indicate that the participants engaged in thoughts recruiting visual imagery during the rest intervals. It is accordingly crucial to use at least one other baseline that matches the experimental problems in everything except the cognitive process of interest. Another reason for the lack of activation in the contrast with rest intervals might be the fixation cross that was displayed throughout the entire experiment to minimize eye movements (see Methods). Possibly, this visual input was more strongly attended to during the rest interval, thus covering other neural activities in the contrast of visual problems and the rest interval.

One interpretation of our results is that they reflect the well-known distinction between the “what” and the “where” pathways in visual processing and working memory. This distinction has been supported by neuroimaging studies of transformation tasks such as mental rotation (e.g., Barnes et al., 2000; Belger et al., 1998; Alivisatos & Petrides, 1997; Cohen et al., 1996; Newcombe et al., 1987; Ungerleider & Mishkin, 1982; Ungerleider, Courtney, & Haxby, 1998). It is also consistent with Kosslyn's hypothesis that mental imagery relies on two different sorts of processes—one visual and one spatial (e.g., Kosslyn, 1994).

Our own interpretation of the results is that they reflect two different sorts of mental representation. Vision yields

a series of representations: Early processes construct images, that is, 2-D representations of intensities, textures, groupings, and so forth, but later vision yields a 3-D model of what is where (Marr, 1982). The phenomena of visual imagery sometimes depend on the manipulations of 2-D images, but sometimes on the manipulation of 3-D models. The time taken to judge whether two pictures are of the same object increases linearly with the angular difference between the orientations of the object in the two pictures (Shepard & Metzler, 1971). This result holds for rotations in the picture plane, but it also holds for rotations in depth. As Metzler and Shepard (1982, p. 45) wrote: “These results seem to be consistent with the notion that . . . subjects were performing their mental operations upon internal representations that were more analogous to three-dimensional objects portrayed in the two-dimensional pictures than to the two-dimensional pictures actually presented.” In other words, the participants were rotating models, not images. Models that may be spatial in form (Johnson-Laird, 1998; Kosslyn, 1994, p. 324) appear to underlie all sorts of reasoning, but the contents of an inference may also invoke visual images that are not essential to the process of reasoning and can impede the process. In a reasoning task, individuals develop facility in constructing mental models from all sorts of relations. Such models will be spatial in form for visuospatial and spatial relations, and, as long-standing evidence suggests, control relations are also likely to elicit spatial models (see, e.g., Johnson-Laird, 1998; De Soto et al., 1965). However, in the case of visual relations, such as “The dog is dirtier than the cat,” reasoners cannot suppress a spontaneous visual image of the appearance of the animals: They are both dirty. Its construction calls for additional activity in visual association cortex. It retards the construction of a mental model that represents the relative degrees of dirtiness, and this model is essential for the inferential process. Hence, the construction of the image slows the process of reasoning.

Is there an alternative explanation for our results? An obvious alternative is that there is some other aspect of the four kinds of relations that distinguishes between the visual relations and the remaining relations, apart from the tendency of the visual relations to elicit images. Number of syllables, syntactic structure, and frequency of usage, all fail to yield the appropriate distinction in both English and German. Another possibility is that some differences in meaning between the various sorts of relations and in their retrieval from long-term memory somehow account for the phenomena. However, as previous studies of such effects have implicated, not visual cortex, but medio-temporal structures, such as the hippocampal formation (see Gabrieli, 1998; Squire & Zola, 1996), and areas in prefrontal and anterior cingulate cortices (Tulving & Craik, 2000) are activated by retrieval from long-term memory. Another possibility is that the relations differ in their discriminability. It may be harder to discriminate between

entities related, say, according to how clean they are, than between entities related according to how smart they are. This hypothesis seems improbable because it is again unable to explain the differences in activity in secondary visual cortex.

Our results have implications for other theories of reasoning and of the role of working memory. A major alternative to the model theory is that reasoners rely on formal rules of inference akin to those of formal logic, and that inference is a process of proof in which the rules are applied to mental sentences (e.g., Braine & O'Brien, 1998; Rips, 1994; but cf. Stenning & Oberlander, 1995). The formal rules govern sentential connectives such as "if" and quantifiers such as "any," and they can account for relational inferences when they are supplemented with axioms governing transitivity such as: For any x , y , and z , if x is taller than y and y is taller than z , then x is taller than z . The formal rule theories accordingly imply that reasoning is a linguistic and syntactic process, and so it should depend on regions in left hemisphere. The model theory, however, postulates that a major component of reasoning is not verbal, and so "the theory predicts that the right cerebral hemisphere should play a significant role in reasoning" (Johnson-Laird, 1995, p. 1005). Wharton and Grafman (1998) have argued that the effects of brain damage on reasoning corroborate formal rule theories and run counter to the model theory. However, our results, in common with other studies (e.g., Kroger, Cohen, & Johnson-Laird, submitted), cast doubt on this analysis. Linguistic areas underlie the comprehension of the premises. Reasoning itself, as the model theory predicts, also implicates parietal regions thought to mediate spatial representations, including regions in right hemisphere. Such activity is inexplicable according to formal rule theories of reasoning.

Reasoning yields activity in DLPFC, pointing to additional executive processes in working memory (Duncan & Owen, 2000; Cohen, Botvinick, & Carter, 2000; Carter, Botvinick, & Cohen, 1999; Smith & Jonides, 1999; Bush et al., 1998). Petrides (1995, p. 85) argues that this region is typically implicated "when several pieces of information in working memory need to be monitored and manipulated." This view has its supporters (see D'Esposito, 2001; Owen, 2000; Petrides, 2000; D'Esposito et al., 1998) and its critics (Levy & Goldman-Rakic, 2000; Mecklinger, Bosch, Gruenewald, Bentin, & von Cramon, 2000; Courtney, Ungerleider, Keil, & Haxby, 1996). A related possibility is that the region underlies the integration of multiple relations. Waltz et al. (1999) showed that patients with damage to prefrontal cortex were strongly impaired in any sort of reasoning calling for the integration of relations, whereas they performed normally in episodic and semantic memory tasks. It seems reasonable to conclude that activation in this region reflects the manipulation or integration of mental representations.

Conclusions

The cortical activity evoked by relational reasoning depends on the nature of the relations. Visual relations, such as "cleaner" or "dirtier than," evoke visual images, whereas other relations do not. All relations, however, lead to the construction of models that underlie the inferential process. These models are spatial in form for spatial relations, such as "north" or "south of," and for visuospatial relations, such as "above" or "below." They may also be spatial for control relations, such as "better" or "worse than." This view is consistent both with the finding that visual relations slow the process of reasoning and activate regions in secondary visual cortex, and with the finding that all relations activate regions in parietal cortex underlying spatial representations. The phenomena are consistent with the theory of mental models, but present difficulties for theories of reasoning based on formal rules of inference. A further corollary is that visual imagery is not a mere epiphenomenon playing no causal role in reasoning (e.g., Pylyshyn, 1981, in press). It can be a nuisance because it impedes reasoning.

METHODS

Norming Studies

Because they are essential for this article, we describe the gist of the studies carried out by Knauff and Johnson-Laird (2002). In the norming studies, we selected 15 pairs of relational terms (a relation and its converse) that might be instances of the different sorts of relation, including such pairs as: "cleaner–dirtier," "uglier–prettier," "heavier–lighter," and "smarter–dumber." We formed 30 assertions using these relations, such as "The cat is above the dog" and "The cat is smarter than the dog." Ten student volunteers at Princeton University used two separate scales to rate the ease of forming visual images and the ease of envisaging spatial layouts for each of the assertions. The two scales had seven points, ranging from "very easy" to "very difficult." The ratings reliably discriminated between the ease of envisaging images and spatial layouts, but they did not yield any spatial relations, that is, those that are easy to envisage spatially but not easy to visualize. We therefore carried out a second norming study using a different procedure. The participants rated each relation on a single bipolar seven-point scale, ranging from ease of evoking a "visual image" at one end of the scale to ease of evoking a "spatial layout" at the other end of the scale. The instructions stated that a visual image is a vivid representation that can include people, objects, colors, and shapes, and that it is similar to a real perception. They also stated that a spatial layout is more abstract and represents an arrangement in terms of a scale or array. We tested 20 students from Freiburg University with a set of 35 relations. This study was successful in identifying all four sorts of relations, including spatial relations.

sorts of relations (visual, visuospatial, and control), yielding a total of 24 inferences. The inferences were presented in a different random order to each participant. The procedure was identical to that of the first experiment, and there were four practice trials before the experiment proper.

In Experiment 3, 24 students of Freiburg University acted as their own controls and carried out 16 three- and 16 four-term series inferences that were identical in form to those in the first experiment. The inferences were based on four kinds of relations: spatial, visuospatial, visual, and control relations. The spatial relations were “ancestor of” and “descendant of” and “further north” and “further south” (all single words in German). There were two valid and two invalid inferences of each of the four sorts in both the three- and four-term series inferences (a total of 32 inferences). These inferences were presented in a different random order to each participant. The procedure was identical to that of the other experiments.

Imaging Study

Participants

We tested 12 male right-handed German native speakers between 21 and 35 years with normal or corrected-to-normal vision, normal hearing, and with no tendency to suffer from claustrophobia (mean age 23.7 years). These volunteers had no magnetic metal in their bodies, no history of neurological or psychiatric disorder, and no history of significant drug abuse. All of them gave their informed consent to take part in the study. As a result of uncorrectable head movements, one participant was excluded from the analysis.

Materials and Experimental Design

The materials were based on the previous behavioral studies (Knauff & Johnson-Laird, 2000, 2002). The participants acted as their own controls and evaluated eight inferences based on each of the four types of relation: visuospatial, visual, spatial, and control relations (see Table 5). All the inferences used the same nouns (“dog,” “cat,” and “ape”) to minimize differences arising from the entities in the problems, for example:

The dog is cleaner than the cat.

The ape is dirtier than the cat.

Does it follow:

The dog is cleaner than the ape?

The participants were told to evaluate whether the conclusion followed from the premises. In the example, “cleaner” and “dirtier” are used once in each premise, and “cleaner” occurs in the conclusion. But, in the experiment as a whole, each relation and its converse occurred equally often in each premise and in the conclusion.

The experiment used a blocked design in which each block contained two problems. The two problems in one block used two different relations of the same sort (visual, spatial, visuospatial, and control). Half of the problems were valid, and half were invalid. The allocation of problems to a block was made at random, and it remained constant throughout the experiment for all participants. The 32 problems were presented in four separate runs wherein each contained four blocks. In this way, we controlled for effects of learning and fatigue and for fluctuations in the scanner’s sensitivity over runs. The order of blocks within each run for each participant was random.

In each block, two problems were presented one after the other. Between each sentence, there was a silent break of 5 sec during which scanning was carried out: The hemodynamic response was thus sampled in the absence of task-related auditory input. A rest interval of similar length was included between the blocks throughout the experiment. This rest interval differed from the silent breaks only because a small fixation cross was presented to minimize eye movements. The cross remained visible on the screen for the duration of the rest interval and through the presentation of each problem. The participants responded to the problems by pressing a “yes” or a “no” button during the response interval after the presentation of each conclusion. To prepare the participants for the start of the next problem, the cross on the screen disappeared one second before the end of the response interval and, where relevant, for one second before the end of the rest interval.

Procedures and Scanning Techniques

Prior to the neuroimaging study, the participants completed a computerized training experiment using the identical conditions, problems, and times as in the neuroimaging experiment. Its aim was to reduce the effects of novelty or of the development of new strategies during scanning. After this training, the participants were given the instructions for the subsequent experiment, and they were placed in the scanner (1.5 T Siemens VISION). Their head position was fixed in the head coil, and a mirror system was placed on the coil so that they could see a projection screen mounted on the rear of the scanner bore. During the experiment, the fixation cross was projected onto this screen using a video beamer. The reasoning problems were presented over the headphones, and the participants responded to them by pressing two buttons of a MR-compatible response box with their right index and middle finger (the “yes” and “no” responses, respectively). The responses were recorded by the program, and the response times were automatically calculated from the presentation of the conclusion until the response. The presentation of each block of two problems was synchronized with the TTL

pulse emitted by the scanner at the start of the collection of each functional image.

Functional images were collected in a gradient-recalled echo-planar imaging (EPI) sequence, allowing the sampling of 24 parallel slices effectively covering the whole brain (TR 10.2 sec, TE 66 msec, fatsat, FA 90°, FOV 256 × 256 mm, 2 × 2 mm in-plane resolution, 4-mm slice thickness). Two rest scans were collected in the beginning of each run in order for T1 effects to stabilize. These images were discarded from further analysis. At the end of the experiment, a functional EPI image was acquired with 40 slices and a sagittal T1-weighted magnetization-prepared rapid-acquisition gradient-echo (MP-RAGE) image of the entire brain (160 slices, TR 40 msec, TE 6 msec, FA 40°, FOV 256 × 256 mm, voxel size 1 × 1 × 1 mm). This image was used for coregistration and normalization during image preprocessing.

Data Analysis

All images were preprocessed and analyzed with the Statistical Parametric Mapping 99 software (SPM99). Functional and anatomical images were reoriented so that the anterior commissure corresponded to the origin of the 3-D standard coordinate system used in SPM99. All coordinates used to describe the results of the experiment corresponded to the system originally introduced by Talairach and Tournoux (1988). The functional images collected during a run were subsequently realigned and corrected for motion in the scanner with the two-step realignment procedure embedded in SPM99 (Friston, Williams, Howard, Frackowiak, & Turner, 1996). The translation and rotation corrections did not exceed 2.9 mm and 2.5° for any run. Each participant's anatomical image series was subsequently coregistered with the 40-slice EPI image volume collected at the end of the experiment (cf. Ashburner & Friston, 1997). The mean images of the runs and all other functional images were then coregistered with the 40-slice EPI image volume. The parameters for spatial normalization (Friston, Ashburner, et al., 1995) were determined from the anatomical image volumes collected from each participant, and they were applied to the participant's functional image volumes. A new set of normalized functional images was created for which the original spatial resolution of the EPI volumes was interpolated from 2 × 2 × 4 mm to an isotropic voxel size of 3 mm³. The realigned normalized images were finally smoothed with a 6-mm full width half maximum Gaussian kernel.

The hemodynamic response to the experimental cycle was modeled with a fixed-response box-car function, which was convolved with the canonical hemodynamic response function to model the delayed onset, early peak, and late undershoot of the hemodynamic response (Veltman & Hutton, 2001). Low-frequency confounds were excluded from the model with a high-pass filter of twice the length of the experimental run. Variations in

global signal intensity were not removed, because of the danger of producing spurious local changes in the direction opposite to any change in the global signal. For each experimental run, the function contained the appropriately placed models of the hemodynamic response to the four sorts of problems. The realignment parameters for each run were included in the model as covariates to correct for the effects of head movement on the hemodynamic response. In the special case of colinearity of realignment parameter sets (intercorrelation $p > .80$), we included only the set showing lower correlations with the remaining four parameter sets.

The inferential task was complicated, it depended on many different cognitive processes, and the block design relied on an extended time scale. Hence, we expected activation differences for all contrasts to be widely spread rather than clearly focused. We therefore made analyses on all contrasts at the level of clusters (Poline, Holmes, Worsley, & Friston, 1997; Friston, Holmes, Poline, Price, & Frith, 1995). SPMs of all contrasts were kept at a threshold of $z = 3.1$, corresponding to $p < .001$, and we identified all clusters that were significant in terms of their size ($p < .05$ corrected for multiple comparisons across the whole brain, resulting in a cluster size of 15 voxels).

For the more fine-grained second comparison described in Results, the four contrasts (visual, visuo-spatial, spatial, and control inferences vs. the rest interval) were examined with a form of conjunction analysis. For this purpose, a new contrast was specified between the images acquired during all experimental conditions and those acquired during the rest intervals. This contrast was inclusively masked with the contrast images calculated for the four single contrasts, which were initially held at a threshold of $p = .05$ uncorrected for multiple comparisons. This analysis displayed only those voxels associated with significantly higher signal intensity during reasoning with all sorts of relation as compared to the rest intervals, and it did not display any voxels significant for reasoning with only one sort of relation. The analysis is not, strictly speaking, a conjunction analysis, for which an intersection SPM(T) is calculated for the single contrasts to perform statistical inference at the voxel level only (Friston, 1997).

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The data reported in this experiment have been deposited in The fMRI Data Center (<http://www.fmridc.org>). The accession number is 2-2002-113BQ.

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