Anatomical and functional subdivision within the primate lateral prefrontal cortex

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It has been suggested that there is specialization of function within the lateral prefrontal cortex; it has been argued that the ventral and dorsal divisions of the prefrontal cortex are concerned with the processing of form and spatial information, respectively. The evidence for such a division of function is reviewed. According to an alternative view, the dorsal prefrontal cortex is better thought of as playing a high-level role within the context of the motor system; this is consistent with the fact that it is this part of the prefrontal cortex that is connected with lateral and medial premotor areas, the parietal cortex, and the cerebellum and that it is this part of the prefrontal cortex that is concerned with action selection and attention to action. The ventral prefrontal cortex, on the other hand, has widespread connections with sensory cortices and may be essential for stimulus selection and sensory attention; this may underlie its role in arbitrary or conditional associations involving diverse stimuli.

Every year physiologists and anatomists repack the primate temporal and parietal lobes into ever smaller divisions. Each region has its own distinctive pattern of neuronal activity and anatomical connections that allows it to make its own individual contribution to the cognitive life of the brain's owner. The frontal lobes stand as something of an affront to the subdividers' project, because there is so little agreement about what the basic component processes of the frontal lobes might be (Roberts, Robbins, & Weiskrantz, 1998).

It is less contentious to suggest that we can draw basic distinctions between the lateral and the ventral surfaces of the prefrontal cortex (see, e.g., Dias, Robbins, & Roberts, 1996; Roberts, 1996). Perhaps the most influential theory of lateral prefrontal function has concerned its role in holding (Goldman-Rakic, 1987, 1998) and manipulating (Petrides, 1998) information in working memory. In people, working memory deficits after lateral prefrontal lesions can be distinguished from deficits in the affective control of behavior caused by lesions of the ventromedial prefrontal cortex (Bechara, Damasio, Tranel, & Anderson, 1998). Neuroimaging studies also suggest a similar role for the human ventromedial prefrontal cortex (Elliott, Frith, & Dolan, 1997, 1999). In the monkey, a high proportion of ventromedial prefrontal cells are concerned with learning about potential rewards and punishments (Rolls, 1999), and lesions here disrupt a monkey's ability to learn which one of a pair of objects is rewarded (Baylis & Gaffan, 1991). What is not so clear, however, is whether there are any functional subdivisions to be found within the large expanse of the lateral prefrontal cortex.

Working Memory and Regional Specialization Within the Lateral Prefrontal Cortex

Any reference to working memory must be prefaced by acknowledgement of the confusion that has surrounded the term. The following discussion of working memory will be restricted to the more limited sense of retention of information through short delays in order to guide subsequent action (Goldman-Rakic, 1987). This hypothesis is distinct from the more complicated set of processes that include "executive control," manipulation of information, the control of attention, and short-term memory that have been subsumed under the heading of working memory by authors such as Baddeley and Hitch (1974).

According to Goldman-Rakic's (1998) domain-specific theory of prefrontal function, different regions of the lateral prefrontal cortex are concerned with holding different types of information. Whereas the ventral lateral prefrontal cortex (VLPFC) is concerned with holding information about forms, the dorsolateral prefrontal cortex (DLPFC) is concerned with holding information about spatial locations. Two principle lines of evidence have been adduced in support of this model.

First, it has been argued that the DLPFC and the VLPFC are distinguished by their anatomical connections with the parietal and the temporal lobes, respectively (Petrides & Pandya, 1984). An influential account of parietal and temporal lobe specialization (Ungerleider & Mishkin, 1982) maintains that these regions are relatively specialized for processing spatial and object-related information, respectively. It was therefore argued that the specializations of the parietal and temporal lobes might be carried forward into the DLPFC and the VLPFC. The attractively simple picture of temporal--VLPFC and parietal--DLPFC...
systems should be revised by demonstrations that the anatomical connections are not so clearly segregated; there are some connections between some more posterior parts of the VLPFC and the parietal lobe (Cavada & Goldman-Rakic, 1989; Schall, Morel, King, & Bullier, 1995). Nevertheless, despite these equivocations, hypotheses of temporal/parietal specialization can still inform hypotheses of VLPFC/DLPFC specialization.

The second line of evidence is based on claims that monkey lesion and cell recording studies and human neuroimaging studies have demonstrated dorsal and ventral specializations for processing spatial and form information, respectively. The human neuroimaging literature will be briefly reviewed before considering the results of cell recording experiments performed with monkeys. Rushworth and Owen (Owen 1997; Rushworth, Nixon, Eacott, & Passingham, 1997; Rushworth & Owen, 1998) have reviewed this evidence and have argued that it is not always persuasive. For example, Figure 1 summarizes activations recorded in human neuroimaging studies of object/form or spatial working memory tasks published before 1998. It is clear that there is no dorsal/ventral distinction between spatial and form tasks. The only evidence that form and spatial information processing is segregated within the prefrontal cortex is that there may be a rostral/caudal division within the VLPFC (Figure 1). Possibly related rostral/caudal transitions have recently been observed in macaque single-unit cell recordings (White & Wise, 1999). Rainer, Asaad, and Miller (1998a) have also emphasized a more prevalent spatial signal in the posterior prefrontal cortex. Human neuroimaging studies have shown that there may also be a dorsal region just in front of the frontal eye fields, outside of what is classically considered the DLPFC, that is specialized just for spatial working memory tasks (Courtney, Petit, Maisog, Ungerleider, & Haxby, 1998). What may be an analogous part of the macaque brain, in the anterior part of area 8, has also been shown to contain cells with delay activity (Chafee & Goldman-Rakic, 1998).

On turning to the monkey single-unit recordings studies that have examined DLPFC/VLPFC specializations for form and spatial processing, it is clear that the experiments have produced inconsistent results that are difficult to reconcile. On the basis of single-unit recording studies, it has been claimed that there is evidence both for (O'Scalaidhe, Wilson, & Goldman-Rakic, 1997, 1999; Wilson, O'Scalaidhe, & Goldman-Rakic, 1993) and against (Fuster, Bauer, & Jervey, 1982; Rao, Rainer, & Miller, 1997) anatomical segregation within the prefrontal cortex. A similar dichotomy of opinion exists when one considers monkey lesion studies (Levy & Goldman-Rakic, 1999; Petrides, 1995; Rushworth, Nixon, et al., 1997). The debate has been reviewed at length before (Rushworth & Owen, 1998), and rather than detaining ourselves with it further, it may be better to summarize: There may be some segregation of function between the dorsal and the ventral prefrontal cortices, but if there is, its basis and guiding principle remain unclear. The next section considers an alternative proposal for functional specialization within the lateral frontal cortex.

A Reevaluation of the Importance of the Lateral Prefrontal Cortices' Anatomical Connections with the Parietal and Temporal Lobes

The inference from the lateral prefrontal cortex's connections with the temporal and parietal lobes to domain-specific processing within the prefrontal cortex is itself based on a particular conception of parietal and temporal lobe function (Ungerleider & Mishkin, 1982). Ungerleider and Mishkin argued that the parietal and temporal lobes were the destinations of two streams of visual-processing areas. It was claimed that the dorsal stream, which projects to the parietal cortex, was concerned with visuospatial processing, whereas the ventral stream, which projects to the temporal lobe, was concerned with visual form discrimination. The theory had broad explanatory power and, as a result, has been one of the most influential in cognitive neuroscience. Nevertheless, it has been challenged. For the present argument, what is at issue is not
whether it is perhaps more realistic to consider whether the two streams are further subdivided (Van Essen & De Yoe, 1995; Young, 1992) or whether they are not strictly serial (Hupe, James, Payne, & Lomber, 1998). Instead, what is at issue is the precise nature of the functional specializations of the parietal and temporal lobe areas. Goodale and Milner (1992; Milner & Goodale, 1995) argued that the functional distinction between the parietal and the temporal lobes was not to be conceived of in terms of specializations for processing different kinds of visual information but in terms of specializations for processing visual information for different purposes; they argued that the ventral and the dorsal visual streams were specialized for using visual information for object identification and for movement guidance, respectively.

At first, the theoretical difference seems subtle, but it has important ramifications. Some of the visually responsive areas of the parietal cortex, such as the lateral intraparietal area (LIP) are now known to contain cells with activity rates that are modulated by visual shapes (Sakata et al., 1999; Sereno & Maunsell, 1998). Such a finding is difficult for a theory based on the distinction between form and spatial vision (Ungerleider & Mishkin, 1982), but it can be accommodated by a theory based on a distinction between vision for identification and vision for action (Milner & Goodale, 1995). As Milner and Goodale point out, visual forms are not just important for helping us to distinguish between objects. They also determine the manner in which movements are made; grasping and other hand movements are shaped by the visual form of the objects that are targeted. Moreover, the ability of both people and monkeys to use visual form information to preshape hand movements is lost after parietal lesions (Perenin & Vighetto, 1988; Rushworth, Nixon, & Passingham, 1997).

If the temporal and parietal lobes are not defined by the type of information that they process, it seems unlikely that different parts of the prefrontal cortex, with which they are interconnected, will be domain-specific processing areas for form and space. Instead, the relative specializations of the VLPFC and the DLPFC are more likely to reflect the temporal and parietal lobes’ respective concerns of stimulus identification and action guidance and selection. In order to achieve these goals, visual information that is either form related or spatial will be processed by either lateral prefrontal region.

In this section, it was considered whether the VLPFC and the DLPFC were specialized for form and spatial working memory. If, however, the VLPFC and the DLPFC are preferentially interconnected with the temporal and the parietal lobes, respectively, then, according to Milner and Goodale’s (1995) theory, we should expect the VLPFC and the DLPFC to be preferentially involved in visual identification and movement, respectively. In the next section, it will be considered whether a DLPFC specialization for movement-related processes and a VLPFC specialization for visual identification processes is consistent with our current knowledge of the other connections of the prefrontal cortex of the monkey.

Other Anatomical Connections of the Prefrontal Cortex

When we move beyond just the temporal and parietal lobes and consider a wider view of the lateral prefrontal cortex’s connections, it is clear that the most distinguishing feature of the DLPFC is not just that it is connected with the dorsal visual stream, but that it is connected with motor regions of the brain. Although the DLPFC does not project directly to the primary motor cortex (M1), when we consider its projections to the premotor areas (on both the lateral and the medial surface) and to the cerebellum, a clear pattern emerges: The DLPFC’s connections with the parietal lobes are just one of the many that the DLPFC has with motor-related brain structures.

The sulcus principalis region of the macaque DLPFC projects to three motor areas on the medial surface of the frontal lobe (Bates & Goldman-Rakic, 1993; Lu, Preston, & Strick, 1994): the presupplementary motor area (pre-SMA) and the most anterior parts of the supplementary motor area (SMA), the rostral cingulate motor area (CMAr) and the ventral cingulate motor area (CMAv). Although the pre-SMA itself has no direct connections with the spinal cord, the adjacent SMA has many, and it is connected with M1 (Dum & Strick, 1996; He, Dum, & Strick, 1995; Lu et al., 1994). Parts of the cingulate motor areas, CMAr and CMAV, have connections with anterior gyral parts of M1 and with the spinal cord (Dum & Strick, 1993, 1996; He et al., 1995; Lu et al., 1994). The VLPFC shares none of these connections (Morecraft & Van Hoesen, 1993; Van Hoesen & Morecraft, 1993; Van Hoesen, Morecraft, & Vogt, 1993).

The sulcus principalis region of the macaque DLPFC also projects to the motor areas on the lateral surface of the frontal lobe: the dorsal premotor cortex (PMd) and the ventral premotor cortex (PMv). Again both these regions, or at least parts of them, have connections with M1 and the spinal cord (Dum & Strick, 1991; Galea & Darian-Smith, 1994; He, Dum, & Strick, 1993). The VLPFC has no connection with the PMd. The issue of whether the VLPFC has connections with the PMv is not clear. Pandya and Vierterian’s (1998) diagrams indicate that there may be some VLPFC-to-PMv connections, but other studies suggest that the origin of the lateral prefrontal connections with the PMv is restricted to the principalis region of the DLPFC (Luppino, Govoni, & Matelli, 1998; Matelli, Camarda, Glickstein, & Rizzolatti, 1986). Carmichael and Price (1995) claimed that the VLPFC does not have direct PMv connections, but they did find evidence of connections from the adjacent ventromedial frontal area (12m) to the PMv.

The DLPFC is also interconnected with subcortical motor structures. It projects to the superior colliculi (Goldman & Nauta, 1976), which are concerned principally with eye movements, but may also play a role in directing limb
Figure 2. Summary diagram of the traditional conception of the anatomical connections of the lateral prefrontal cortex with the temporal and parietal lobes. The dorsolateral prefrontal cortex (DLPFC) is connected with the dorsal visual stream and the parietal cortex. The ventrolateral prefrontal cortex (VLPFC) is connected with the ventral visual stream and the temporal lobe.

movements (Stuphorn, Hoffmann, & Miller, 1999; Werner, Hoffmann, & Dannenberg, 1997). Finally, it is just the DLPFC, and not the VLPFC, that projects to the cerebellum via the pons (Middleton & Strick, 1998; Schmahmann & Pandya, 1997).

Instead of motor connections, the VLPFC has access to a range of highly processed sensory information. Most noticeable is the manner in which the VLPFC is interconnected with the visual and auditory modalities; these are probably the most important senses for object identification in primates. There are strong connections between the anterior VLPFC and the inferotemporal visual cortex (Carmichael & Price, 1995; Pandya & Yeterian, 1998; Webster, Bachevalier, & Ungerleider, 1994). A slightly more posterior division of the VLPFC is interconnected with the auditory superior temporal gyrus (Pandya & Yeterian, 1998). It is also the VLPFC that is interconnected with the perirhinal cortices of the temporal lobe, which have a particularly important role in representing knowledge about objects (Murray & Bussey, 1999) in several modalities (Suzuki, Zola-Morgan, Squire, & Amaral, 1993).

In conclusion, a reconsideration of the specializations of dorsal and ventral visual stream areas (Milner & Goodale, 1995) suggests that the DLPFC and the VLPFC are specialized for motor- and sensory-related processes, respectively. A broader consideration of the anatomical connections of the two areas is consistent with this hypothesis; the DLPFC is interconnected with many pre-motor areas and subcortical motor structures, such as the cerebellum, whereas the VLPFC has access to information regarding object identity from several modalities via its connections with the temporal lobe. The anatomical connections of the lateral prefrontal cortex are summarized in Figures 2 and 3.

Sensory Versus Motor Learning in the Lateral Frontal Lobes

The last section reviewed the anatomical evidence for VLPFC’s and DLPFC’s interconnections with sensory- and motor-related brain regions, respectively. This section reviews neuroimaging evidence as to whether the VLPFC and the DLPFC are relatively specialized for sensory- or motor-related processes.

It has been argued that the prefrontal cortex is specialized for only certain types of mnemonic task, such as working memory tasks, in which various items of information must be stored for short periods of time (Goldman-Rakic, 1998). In the case of the VLPFC, however, it is clear that lesions impair the learning of tasks with no delay element while having little effect on monkeys’ abilities to hold information for short periods of time (Murray, Bussey, & Wise, in press; Parker & Gaffan, 1998; Passingham, 1972; Rushworth, Nixon, et al., 1997). Some authors have gone so far as to suggest that the prefrontal cortex is best considered as a “last-ditch general learning mechanism” for learning tasks that cannot be managed by the various, more specialized learning mechanisms of the medial temporal lobe (Gaffan, 1994). This perspective suggests that it might be useful to consider not whether lateral frontal regions are specialized for spatial or object-based working memory (Figure 1), but whether they are specialized for learning difficult sensory or motor tasks (Figure 4).
In several human neuroimaging studies, changes in brain activation were investigated while subjects learned sequences of movements (Grafton, Hazeltine, & Ivry, 1995; Hazeltine, Grafton, & Ivry, 1997; Honda et al., 1998; Jueptner et al., 1997; Sakai et al., 1998; Toni, Schluter, Josephs, Friston, & Passingham, 1999), and all recorded activation in the DLPFC in the region of the middle frontal gyrus. A ventral prefrontal peak was also recorded in one study (Toni et al., 1999). There is a comparable dearth of neuroimaging studies of perceptual learning, and some of those that exist did not include any scanning of the frontal lobes (Poldrack, Desmond, Glover, & Gabrieli, 1998) but, instead, concentrated on the more posterior visual areas that are known to play a role in visual learning (Walsh, Ashbridge, & Cowey, 1998). Two perceptual-learning studies that did scan the prefrontal cortex, however, both found a number of activation peaks in the VLPFC, but just one activation change in the DLPFC (Pettersson, Elfgren, & Ingvar, 1999; Schiltz et al., 1999). In conclusion, it is mainly the DLPFC that is recruited in motor-learning tasks but the VLPFC that is recruited in perceptual-learning tasks (Figure 4).

Attention in the Lateral Frontal Cortex

Although it is possible to use perceptual- and motor-learning tasks to distinguish the functions of the VLPFC and the DLPFC, this does not mean that these regions are only important during the learning of new tasks. It seems as if it is only necessary to require subjects to attend to their movements for activation to appear in the DLPFC. Jueptner et al. (1997) asked their subjects to perform a previously learned motor sequence in one of two ways. In one case, subjects were simply asked to perform the sequence, but in a second condition, they were asked to pay attention to what they were doing. There was significantly more activity in the DLPFC when the movements were attended. Other studies in which subjects pay attention to even single, simple finger movements also result in activation of the DLPFC (Krams, Rushworth, Deiber, Frackowiak, & Passingham, 1998; Rushworth, Krams, & Passingham, in press). In monkeys, the activity of DLPFC cells is modulated during the preparation of a response (Fuster, 1995; Hasegawa, Sawaguchi, & Kubota, 1998).

It might be argued that when subjects are attending to actions, they are really just attending to the spatial destinations of the actions. All of the studies described above, however, involve just finger movements, and it is not clear that subjects code such simple movements spatially. The review of studies in Figure 1 confirms that the DLPFC is not distinguished by a special role in processing spatial information. Perhaps the most persuasive demonstration

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**Figure 3.** Summary diagram of the anatomical connections of the lateral prefrontal cortex with the motor association cortex, the cerebellum, and the sensory association cortex. From this broader perspective, it can be seen that the connection of the dorsolateral prefrontal cortex (DLPFC) with the parietal lobe is just one of the many connections between the DLPFC and the motor areas of the brain. In addition, it is clear that the connection of the ventrolateral prefrontal cortex (VLPFC) with the inferior temporal lobe is just one of the connections of the VLPFC with sensory association cortices concerned with object knowledge and identification. Pre-SMA, presupplementary motor area; CMAr, rostral cingulate motor area; CMAv, ventral cingulate motor area; PMd, dorsal premotor cortex; PMv, ventral premotor cortex.
that the DLPFC is not just concerned with the spatial dimension of a movement comes from a monkey lesion experiment. Passingham (1978) taught monkeys to press a single button repeatedly until a light was turned off. The monkeys were then required to reproduce approximately the same number of movements on the same button. DLPFC lesions impaired the monkeys' ability to perform the task even when there was no delay interposed between the end of the first movement sequence and the start of the second movement sequence. The results can be explained if the DLPFC lesions prevented the monkeys from attending to their own movements.

Conversely, it seems as if tasks requiring subjects to attend to the visual features of objects result in more ventral activation in the VLPFC. For example, Konishi et al. (1998) recorded activity in the inferior frontal sulcus as subjects switched the focus of their attention between the color and the form of stimuli. When Sakagami and Niki (1994) trained their monkeys to perform a similar task, they found that the cue-related activity of VLPFC cells depended on which dimension of the cue was currently behaviorally relevant—that is, cells’ activity depended on whether the cue was red or green when color was the relevant dimension but depended on whether the cue was a circle or a cross when cue shape was the relevant dimension or on whether the cue was on the left or the right when cue position was the relevant dimension. Rainer et al. (1998a, 1998b) similarly reported that the visual activity of prefrontal neurons was influenced not just by the identity and location of stimuli presented in a visual array, but also by which stimuli and locations were currently the focus of attention. Moreover, they also reported that the visual activity of some cells was unaffected by the presence or absence or the position of unattended stimuli. Rainer et al. (1998a) provide only a brief account of the anatomical location at which they recorded, but their Figure 1 suggests that the cells that they were recording from were mainly below the principal sulcus in the VLPFC. Face stimuli are very salient stimuli for both monkeys and people, and in this case, it seems that there is less doubt that they are preferentially processed in the VLPFC of both species, even outside the context of any working memory task (Courtney et al., 1998; Haxby et al., 1994; O'Scalaidhe et al., 1997, 1999).

Action Selection in the Lateral Frontal Cortex

The frontal lobes are required when actions are to be selected, particularly when the action must be selected in an unusual or nonroutine situation or when the rules guiding the selection have to be learned. Once again, it is possible to find some evidence for regional segregation within the lateral frontal cortex when we compare two different types of action selection tasks, often referred to as willed action and conditional motor selection tasks.

In the willed action, or free selection, task, subjects are allowed to freely choose any response, often out of a free choice between many responses (Deiber, Ibanez, Sadato, & Hallet, 1996; Deiber et al., 1991; Frith, Friston, Liddle, & Frackowiak, 1991; Hyder et al., 1997; Jahanshahi et al., 1995; Jueptner et al., 1997; Sakai et al., 1998; Toni, Schluter, Josephs, Friston, & Passingham, 1999), but a predominance of ventrolateral prefrontal cortex (VLPFC) activation when subjects are learning perceptual tasks (Petersson, Elfgren, & Ingvar, 1999; Schiltz et al., 1999).
limited predefined set on any trial. In different versions of the task, subjects may make verbal responses or limb movement responses. Because there is some evidence that different prefrontal regions are activated when responses are made in different modalities (Hyder et al., 1997), Figure 5 shows only those activations recorded when subjects are performing free selection tasks in which the forelimb is used to make the response. When subjects are choosing between which finger to move, which way to move a joystick, or when to move, there is activation in the DLPFC (Deiber et al., 1996; Hadland et al., 1999; Toni et al., 1999; Park et al., 1999). In fact, the region of the prefrontal cortex that is critical for conditional task performance (Figure 5) appears to be almost immediately situated between the visual-learning and the motor-learning prefrontal regions identified in the previous section (Figure 4).

The Dorsolateral Prefrontal Cortex and the Dorsomedial Frontal Cortex

Emphasizing the DLPFC's role in motor learning and attention to action helps to distinguish its functional role from that of the VLPFC, but at the same time, it unfortunately highlights the current need to distinguish the functional contributions of the DLPFC and the medial frontal areas, particularly the pre-SMA and the anterior cingulate cortex. One or the other or both of these medial frontal areas are frequently activated by the movement sequence learning tasks (Grafton et al., 1995; Hazeltine et al., 1997; Honda et al., 1998; Jenkins, Brooks, Nixon, Frackowiak, & Passingham, 1994; Jueptner et al., 1997; Sakai et al., 1998; Toni et al., 1999), the attention to action tasks (Jueptner et al., 1997; Kram et al., 1998; Rushworth et al., in press), and the free selection tasks (Deiber, Honda, Ibanez, Sudato, & Hallet, 1999; Deiber et al., 1996; Deiber et al., 1991; Frith et al., 1991; Jueptner et al., 1995; Hallet, 1999; Deiber et al., 1996; Honda et al., 1998; Jenkins, Brooks, Nixon, Frackowiak, & Passingham, 1994; Jueptner et al., 1997; Sakai et al., 1998; Toni et al., 1999) described above. It may often be difficult to decide which of these two areas has been activated in any given study (Crosson et al., 1999). In reviewing 107 PET studies, Koski and Paus (in press) emphasized the co-occurrence of activity in the DLPFC and the anterior cingulate cortex. The distinct functional contributions of the DLPFC and the medial frontal cortex may be better revealed by methods with a finer temporal resolution than most current human neuroimaging methods, such as a single-unit recording and event-related fMRI, or by interference studies involving permanent lesions or transient disruption with transcranial magnetic stimulation (Hadland et al., 1999).

Conclusions

It has been argued that it may not be correct to consider the DLPFC and the VLPFC to have distinct roles in spatial and form-related working memory, respectively
Instead, the DLPFC and the VLPFC can be usefully considered to have high level roles in tasks that demand novel, nonroutine, or attentionally demanding ways of processing motor or visual information. This argument has been based on a reconsideration of the functions of the temporal and parietal lobes (Milner & Goodale, 1995), the distinct anatomical connections of the DLPFC and the VLPFC (Figure 3), and a review of neuroimaging, lesion, and single-cell recording studies of visual and motor learning, visual and motor attention, and different types of action selection tasks (Figures 4 and 5).

In some ways, this conclusion echoes a hypothesis put forward by Passingham (1993), before much of the anatomical and neuroimaging data reviewed above had been published. Passingham (1993, p. 208) argued that the VLPFC was "informed about the external context, that is, about the world," whereas the DLPFC was "concerned with the animal itself, its eyes and limbs."

It should be emphasized that the differential specializations of areas are likely to be relative, rather than absolute, when those areas are anatomically interconnected and are required to work in concert to perform certain cognitive tasks. This is not just the case in the lateral prefrontal cortex (Carmichael & Price, 1996; Pandya & Yeterian, 1998) but also applies in the case of the dorsal and ventral visual streams (Van Essen & De Yoe, 1995).

It is possible that this simple framework can be extended to accommodate other findings concerning the prefrontal cortex's role in more complex tasks. For example, the ability to combine and manipulate movement elements into numerous possible sequences is an important feature of primate motor behavior, and both the human and the monkey DLPFC is important for the learned production of movement sequences (Barone & Joseph, 1989; Grafton et al., 1995; Hazeltine et al., 1997; Honda et al., 1998; Jenkins et al., 1994; Jueptner et al., 1997; Sakai et al., 1998; Toni et al., 1999). It may be that this ability to learn and produce sequences, perhaps originally acquired in the context of motor behavior, has been adapted so that the DLPFC now has a role in combining and manipulating internal representations of items in memory (Owen, 1997; Owen, Doyon, Petrides, & Evans, 1996; Owen, Evans, & Petrides, 1996; Petrides, 1995, 1998).

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