Research report

Dorsolateral prefrontal contributions to human working memory

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Abstract

Although neuroscience has made remarkable progress in understanding the involvement of prefrontal cortex (PFC) in human memory, the necessity of dorsolateral PFC (dlPFC) for key competencies of working memory remains largely unexplored. We therefore studied human brain lesion patients to determine whether dlPFC is necessary for working memory function, administering subtests of the Wechsler Memory Scale, the Wechsler Adult Intelligence Scale, and the N-Back Task to three participant groups: dlPFC lesions (n = 19), non-dlPFC lesions (n = 152), and no brain lesions (n = 54). DlPFC damage was associated with deficits in the manipulation of verbal and spatial knowledge, with left dlPFC necessary for manipulating information in working memory and right dlPFC critical for manipulating information in a broader range of reasoning contexts. Our findings elucidate the architecture of working memory, providing key neuropsychological evidence for the necessity of dlPFC in the manipulation of verbal and spatial knowledge.

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1. Introduction

Working memory comprises a system for maintaining, monitoring and manipulating information in short-term memory, providing the interface between perception, long-term memory and action that enables goal-directed behavior (Baddeley, 1998; Baddeley and Petrides, 1996). Although cognitive neuroscience has made remarkable progress in understanding the involvement of the prefrontal cortex (PFC) in human memory, fundamental questions remain regarding the functional organization of the PFC with respect to working memory. One unresolved issue concerns whether subregions within the lateral PFC mediate functionally distinct processes or instead serve a common role in working memory. Anatomically, the lateral PFC consists of multiple subregions that differ in cytoarchitecture and connectivity (Petrides et al., 1996).
2012), raising the possibility that these subregions may guide goal-directed behavior through different mechanisms.

A seminal and longstanding debate in cognitive neuroscience has examined this issue, investigating alternative models for understanding the functional organization of the lateral PFC and its role in working memory. Domain-general models posit that the lateral PFC is functionally organized according to the type of working memory operations engaged, with the dorsolateral PFC (dlPFC) embodying computational mechanisms for monitoring and manipulating items in working memory (Owen et al., 1996; Duncan and Owen, 2000; Miller and Cohen, 2001; Koechlin et al., 2003; Petrides, 2000, 2005; Petrides et al., 2012). Monitoring operations are thought to support the active retention of information in working memory and computational mechanisms for manipulating items are recruited for updating (Petrides, 2000) or selecting between these representations (Rowe et al., 2000). In contrast, domain-specific models posit that the lateral PFC is functionally organized according to the domain of information processed. Advocates of this framework propose that dlPFC is functionally specialized to process visuospatial information in working memory, enabling mental representations of coordinates within the spatial domain (Awh et al., 1995; Butters and Pandya, 1969; Butters et al., 1971, 1972; Courtney et al., 1998, 1996, 1997; Goldman-Rakic, 1995; Levy and Goldman-Rakic, 1999; Smith and Jonides, 1999).

The empirical case advanced in support of each model of dlPFC function has relied primarily upon (1) lesion studies in non-human primates demonstrating reliable deficits in working memory due to unilateral dlPFC lesions (Butters and Pandya, 1969; Butters et al., 1971, 1972; Jacobsen and Nissen, 1937; Levy and Goldman-Rakic, 1999) and (2) functional neuroimaging studies in humans reporting activity within the dlPFC for tests of working memory (for meta-analytic reviews, see (Owen et al., 2005; Wager et al., 2004; Wager and Smith, 2003]). Two key findings from studies of non-human primates performing delayed-response tasks suggest a crucial role for the dlPFC in working memory. First, experimental lesions of the principal sulcus in the dlPFC cause delay-dependent impairments, whereby forgetting increases with the length of the delay (Miller and Orbach, 1972; Bauer and Fuster, 1976; Funahashi et al., 1993). Second, neurophysiological unit recordings from the dlPFC often show persistent, sustained levels of neuronal firing during the retention interval of delayed-response tasks (Funahashi et al., 1989; Fuster and Alexander, 1971; Kubota and Niki, 1971). This sustained activity is thought to provide a bridge between the stimulus cue (e.g., the location of a flash of light) and its contingent response (e.g., a saccade to the remembered location). Such data established a strong link implicating the dlPFC as a crucial node supporting working memory.

Conclusions drawn from these literatures, however, are characterized by the following well-known limitations. First, the precise localization of working memory functions cannot be directly transposed from monkeys to humans due to significant interspecies macroscopic anatomical differences (Petrides et al., 2012). Second, functional neuroimaging (fMRI) studies apply correlational methods and therefore cannot formally demonstrate whether dlPFC is necessary for working memory or instead serves an accessory role (Sarter et al., 1996). As a consequence, the precise localization of working memory function in humans and the contribution of dlPFC to the neural systems underlying working memory remain controversial.

In recent years, lesion studies in humans (Baldo and Dronkers, 2006; D’Esposito and Postle, 1999; D’Esposito et al., 2006; Muller et al., 2002; Pito et al., 1995; Tsuchida and Fellows, 2009; Volle et al., 2008) and repetitive transcranial magnetic stimulation (rTMS) experiments (Hamidi et al., 2009, 2008; Koch et al., 2005; Postle et al., 2006) have provided key evidence to inform the debate. Human lesion and rTMS research are able to overcome the methodological limitations of earlier non-human primate and functional neuroimaging studies by investigating the anatomical localization of working memory functions in the human brain (Rorden and Karnath, 2004) and evaluating the necessity of the dlPFC for specific components of working memory.

Findings from the contemporary literature, however, have been equivocal, with some investigators reporting specific patterns of working memory deficits (Baldo and Dronkers, 2006; Mottaghy et al., 2002; Pito et al., 1995; Tsuchida and Fellows, 2009; Volle et al., 2008) and others failing to observe reliable impairment (D’Esposito and Postle, 1999; D’Esposito et al., 2006; Hamidi et al., 2008; Koch et al., 2005; Muller et al., 2002). Difficulty in interpreting the theoretical significance of these findings has resulted from (1) the often diffuse (rather than focal) lesions observed, (2) the lack of comparison subjects carefully matched for pre- and post-injury performance measures, and (3) the limited scope of working memory functions examined. The absence of such data represents a substantial gap in the understanding of both dlPFC function and the neural substrates of working memory. Here, we characterize key competencies of working memory function in a sample of patients with focal brain lesions involving dlPFC.

2. Materials and methods

2.1. Participant data

We drew brain-injured participants from the Vietnam Head Injury Study (VHIS) registry, which includes American veterans who suffered brain damage from penetrating head injuries in the Vietnam War (n = 199), as well as neurologically healthy Vietnam veterans (n = 54). The VHIS has been organized in three phases. Phase 1 (1967–1970) was the initial enrollment; Phase 2 (1981–1984) included a cognitive evaluation; and Phase 3 (2003–2006) included a more comprehensive evaluation as well as computed tomography (CT) brain imaging. Further details regarding the VHIS participants, including methods for visualizing and quantifying brain lesions, have previously been reported (Barbey et al., 2012a, 2011). Subjects were eligible for the present study if they participated in Phases 2 and 3 evaluations.

To preclude the possibility that impaired performance on working memory and executive function tests could be secondary to deficits in the production and/or comprehension of language, we excluded any participant who had significant impairment on a neuropsychological test of language.
comprehension and production [i.e., defined as performance at least two standard deviations (SDs) below the mean of the neurologically healthy group on the Boston Naming Test]. From the remaining brain-injured veterans we selected those with significant damage to dIPFC (Brodmann’s area 9/46) in the left and/or right hemisphere(s) (dIPFC lesion group; Fig. 1; n = 19). The dIPFC is located on the lateral and dorsal part of the medial convexity of the frontal lobe and comprises BA 9 and 46 and a few transitional areas: 9–8, 9–45, 46–10, and 46–45 (for a detailed description of anatomical boundaries, see Rajkowska and Goldman-Rakic, 1995b, 1995a). In addition, we investigated a comparison group of brain-injured veterans whose damage did not involve dIPFC or the superior parietal lobe, a cortical region necessary for certain aspects of working memory (non-dIPFC lesion group; Supplementary Fig. 1; n = 152; Koenigs et al., 2009). As Supplementary Fig. 1 illustrates, the greatest area of lesion overlap within the non-dIPFC sample entailed the ventral portion of the medial PFC (below the level of the genu of the corpus callosum) and medial portion of the orbital surface (approximately the medial one-third of the orbitofrontal cortex in each hemisphere) as well as the subjacent white matter. Neurologically healthy veterans served as an additional comparison group (no lesion group; n = 54). Demographic and background cognitive function data for the three groups are presented in Table 1. No significant group differences were observed with respect to basic demographic variables (age, sex, years of education), pre- and post-combat measures of verbal IQ and verbal comprehension, and total percent volume loss. All patient groups were therefore well matched with respect to (1) basic demographic variables, (2) pre- and post-combat measures of cognitive function and (3) lesion size.

2.2. Lesion analysis

We acquired CT data during the Phase 3 testing period. Axial CT scans without contrast were acquired at the Bethesda Naval Hospital on a General Electric Medical Systems Light Speed Plus CT scanner in helical mode. We reconstructed the images with an in-plane voxel size of .4 × .4 mm, an overlapping slice thickness of 2.5 mm and a 1-mm slice interval. We determined lesion location and volume from CT images using the Analysis of Brain Lesion (ABLe) software (Makale et al., 2002; Solomon et al., 2007) contained in MEDx v3.44 (Medical Numerics) with enhancements to support the Automated Anatomical Labeling (AAL) atlas (Tzourio-Mazoyer et al., 2002). We applied the AAL atlas of the human brain to obtain neuroanatomical labels for locations in three-dimensional space. For hypotheses about specific brain areas (dIPFC), we defined regions of interest (ROIs) in terms of AAL structures (Tzourio-Mazoyer et al., 2002) and Talairach coordinates. As part of this process, we spatially normalized the CT image of each subject’s brain to a CT template brain image in Montreal Neurological Institute (MNI) space (Collins et al., 1994). We determined the
Table 1 – Demographic and background data.

<table>
<thead>
<tr>
<th>Demographic data</th>
<th>DlPFC</th>
<th>Non-dlPFC</th>
<th>No lesion</th>
<th>ANOVA F value</th>
<th>ANOVA p value</th>
<th>Significant between-group differences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>58.68 (2.75)</td>
<td>58.88 (2.20)</td>
<td>59.52 (3.42)</td>
<td>.91</td>
<td>.41</td>
<td>None</td>
</tr>
<tr>
<td>Sex (% male)</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>n/a</td>
<td>n/a</td>
<td>None</td>
</tr>
<tr>
<td>Years of education</td>
<td>14.74 (2.83)</td>
<td>14.93 (2.53)</td>
<td>15.19 (2.47)</td>
<td>.3</td>
<td>.74</td>
<td>None</td>
</tr>
<tr>
<td>Pre-combat AFQT</td>
<td>54.83 (27.75)</td>
<td>62.40 (24.78)</td>
<td>65.40 (22.91)</td>
<td>1.09</td>
<td>.34</td>
<td>None</td>
</tr>
<tr>
<td>Post-combat AFQT</td>
<td>59.61 (24.20)</td>
<td>67.82 (22.81)</td>
<td>72.34 (22.99)</td>
<td>1.77</td>
<td>.17</td>
<td>None</td>
</tr>
<tr>
<td>Post-combat verbal IQ</td>
<td>102.84 (16.50)</td>
<td>106.89 (12.38)</td>
<td>109.87 (12.38)</td>
<td>2.16</td>
<td>.12</td>
<td>None</td>
</tr>
<tr>
<td>Post-combat verbal comp.</td>
<td>105.00 (17.53)</td>
<td>108.42 (13.60)</td>
<td>109.66 (12.04)</td>
<td>.82</td>
<td>.44</td>
<td>None</td>
</tr>
<tr>
<td>Total percent volume loss (cm³)²</td>
<td>3.11 (1.98)</td>
<td>2.70 (3.59)</td>
<td>n/a</td>
<td>.49</td>
<td>.62</td>
<td>None</td>
</tr>
</tbody>
</table>

Data are presented as means with SDs in parentheses. “Age” refers to age at the time of Phase 3 evaluation. “Sex” refers to the percent of male veterans. “Years of education” refers to the total number of years of education the veterans completed. “Pre-combat AFQT” refers to index scores on the Armed Forces Qualification Test, a battery of tests measuring basic cognitive function at the time of enlistment (pre-injury). “Post-combat AFQT” refers to index scores on the Armed Forces Qualification Test administered at Walter Reed Medical Center after injury. “Post-combat verbal IQ” refers to the Phase 3 verbal IQ index score from the WAIS. “Post-combat verbal comprehension index score” refers to the Phase 3 verbal comprehension index score from the WAIS. Significant between-group differences were determined with the Tukey’s HSD test.

² An independent samples t-test was conducted (rather than an ANOVA) to determine significant between-group differences for the dlPFC and non-dlPFC patient groups. The respective values represent the t score and the associated p value.

2.3. Neuropsychological tests

We administered subtests of the Wechsler Memory Scale, 3rd Edition (WMS III; Wechsler, 1997b), the Wechsler Adult Intelligence Scale, 3rd Edition (WAIS III; Wechsler, 1997b), and an experimental test of working memory, the N-Back Task (Cohen et al., 1997) to investigate the necessity of dlPFC for specific (1) cognitive operations (maintenance, monitoring and manipulation) and (2) modalities of information (verbal and spatial) in working memory. Maintenance operations enable the temporary online retention of information in working memory and are measured by simple retention tasks (e.g., the Digit Span Forward Task). Monitoring refers to the process of deliberately attending to information in working memory and is measured by active retention tasks (the N-Back Task). Manipulating items in working memory refers to the rearrangement and transformation of representations for goal-directed behavior and is measured by tasks that draw upon executive control functions (e.g., the Letter–Number Sequencing Task). The reported neuropsychological data from the WMS III and WAIS III represent standardized scores based on published norms from Wechsler (Wechsler, 1997b). Data for the N-Back Task represent the mean number of errors in each patient group.

2.3.1. Maintenance

We investigated the patient’s ability to maintain information in working memory, administering a verbal/auditory maintenance measure, WAIS III: Digit Span Forward Task, and a non-verbal/spatial maintenance measure, WMS III: Spatial Span Forward Task, which is equivalent to the Corsi Span Task (Kessels et al., 2008). In Digit Span Forward, the patient hears a sequence of digits and attempts to repeat the sequence in order (Wechsler, 1997b). In Spatial Span Forward, the patient watches the examiner tap a sequence of locations on a board and attempts to repeat the tapping sequence in order (Wechsler, 1997b). Together, these tasks provide an assessment of the simple retention of verbal/auditory and non-verbal/spatial representations in working memory.

2.3.2. Monitoring

To examine the patient’s ability to actively monitor information in working memory, we administered the Zero-Back condition of the N-Back Task (Cohen et al., 1997). In this condition, the patient receives a sequence of visually presented letters and indicates whether the letter on the current trial matches a target stimulus. The Zero-Back condition therefore represents a pure measure of monitoring operations, examining the patient’s ability to identify a target stimulus by actively monitoring incoming visual stimuli (Owen et al., 2005).

2.3.3. Cognitive load and processing demands on working memory

We additionally administered the One-, Two-, and Three-Back conditions of the N-Back Task to investigate the recruitment of dlPFC with increasing cognitive load and processing demands on working memory. These conditions support the parametric manipulation of cognitive load, measuring the patient’s ability to determine whether each letter in the series matches the stimulus that occurred either one-, two- or three-trials previously. Successful performance requires that the
patient (1) monitor a series of incoming stimuli, (2) maintain activation of recently processed and potentially relevant items, (3) discard recently processed but irrelevant information, and (4) make comparisons between items in the series to identify a correct match. The One-, Two- and Three-Back conditions of the N-Back Task therefore support an investigation of the dlPFC’s role in working memory with increasing cognitive load and processing demands.

2.3.4. Manipulation

We examined the patient’s ability to manipulate items in working memory, employing two measures of the rearrangement of verbal/auditory information, WMS III: Letter—Number Sequencing and WMS III: Digit Span Backward, and a measure of the manipulation of non-verbal/spatial representations, WMS III Spatial Span Backward. In Letter—Number Sequencing, the patient hears a sequence of alternating digits and letters, and attempts to rearrange the order of each item by repeating the digits in numerical order, followed by the letters in alphabetical order (Wechsler, 1997b). Digit Span Backward (Wechsler, 1997a) and Spatial Span Backward (Wechsler, 1997a) are the same as their forward counterparts, except that the subject attempts to repeat each sequence in reverse order. Together, these measures support an assessment of the manipulation and rearrangement of verbal and spatial representations in working memory.

2.3.5. Reasoning

To investigate whether the dlPFC is necessary for manipulating information in tasks that do not exclusively depend on working memory, we examined manipulation processes in a broader range of verbal and spatial reasoning contexts, administering neuropsychological tests of mental arithmetic, WAIS III: Arithmetic, and visuospatial reasoning, WAIS III: Matrix Reasoning. In Arithmetic the subject hears numerical problems in story format, performs mental arithmetic (i.e., without paper and pencil), and provides a verbal response (Wechsler, 1997b). In Matrix Reasoning, the patient receives pictures of geometric shapes and draws an analogical inference about the missing shape that completes the pattern (Wechsler, 1997b). This task is comparable to Raven’s Progressive Matrices (Raven, 2000). The inclusion of verbal and spatial reasoning tasks complements our analysis of these operations in working memory, supporting an assessment of the contribution of the dlPFC to cognitive operations for manipulating information in a broader range of contexts.

2.3.6. Statistical analyses

We report two main analyses. First, we conducted a one-way analysis of variance (ANOVA) for each neuropsychological measure of working memory and executive function to examine the performance of dlPFC lesion patients ($n = 19$) with respect to non-dlPFC lesion patients ($n = 152$) and neurologically healthy participants ($n = 54$), followed by Tukey’s honestly significant difference (HSD) test to determine significant between-group differences. Second, we conducted a follow-up analysis to investigate the performance of a smaller sample of patients with focal dlPFC lesions, applying non-parametric statistics to test for group effects and for pairwise comparisons.

3. Results

To summarize the results reported in Table 2, no significant group differences in the dlPFC patient sample were observed for neuropsychological tests of working memory maintenance (Digit Span Forward and Spatial Span Forward), monitoring (Zero-Back), or under conditions of increasing cognitive load and processing demands (One-, Two- and Three-Back). However, deficits were observed in the dlPFC patient group for a test of mental arithmetic requiring the manipulation of verbal information (Arithmetic) and approached significance for a working memory test also requiring the manipulation of

<table>
<thead>
<tr>
<th>Cognitive function</th>
<th>Cognitive measure</th>
<th>DlPFC</th>
<th>Non-dlPFC</th>
<th>No lesion</th>
<th>ANOVA F value</th>
<th>ANOVA p value</th>
<th>Significant between-group differences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maintenance</td>
<td>Digit Span Forward</td>
<td>6.05 (1.22)</td>
<td>6.33 (1.20)</td>
<td>6.68 (1.22)</td>
<td>2.47</td>
<td>.09</td>
<td>None</td>
</tr>
<tr>
<td>Maintenance</td>
<td>Spatial Span Forward</td>
<td>9.16 (3.20)</td>
<td>9.52 (3.27)</td>
<td>10.22 (2.69)</td>
<td>1.25</td>
<td>.29</td>
<td>None</td>
</tr>
<tr>
<td>Monitoring: no cognitive load</td>
<td>Zero-Back Errors</td>
<td>1.71 (2.66)</td>
<td>1.26 (1.73)</td>
<td>1.08 (2.33)</td>
<td>.65</td>
<td>.53</td>
<td>None</td>
</tr>
<tr>
<td>Monitoring: low cognitive load</td>
<td>One-Back Errors</td>
<td>3.12 (2.98)</td>
<td>2.98 (2.26)</td>
<td>2.57 (2.48)</td>
<td>.66</td>
<td>.52</td>
<td>None</td>
</tr>
<tr>
<td>Monitoring: medium cognitive load</td>
<td>Two-Back Errors</td>
<td>4.76 (2.02)</td>
<td>4.55 (2.48)</td>
<td>3.85 (2.39)</td>
<td>1.8</td>
<td>.17</td>
<td>None</td>
</tr>
<tr>
<td>Monitoring: high cognitive load</td>
<td>Three-Back Errors</td>
<td>5.65 (2.98)</td>
<td>5.95 (2.66)</td>
<td>4.68 (2.40)</td>
<td>4.39</td>
<td>.01</td>
<td>Non-dlPFC &gt; no lesion*</td>
</tr>
<tr>
<td>Manipulation</td>
<td>Letter—Number Sequencing</td>
<td>9.32 (2.25)</td>
<td>10.08 (2.67)</td>
<td>11.04 (2.66)</td>
<td>3.63</td>
<td>.03</td>
<td>DlPFC &lt; no lesion*</td>
</tr>
<tr>
<td>Manipulation</td>
<td>Digit Span Backward</td>
<td>4.53 (1.35)</td>
<td>4.59 (1.34)</td>
<td>4.96 (1.41)</td>
<td>1.62</td>
<td>.20</td>
<td>None</td>
</tr>
<tr>
<td>Manipulation</td>
<td>Spatial Span Backward</td>
<td>10.58 (3.37)</td>
<td>11.23 (2.89)</td>
<td>12.02 (3.12)</td>
<td>2.09</td>
<td>.13</td>
<td>None</td>
</tr>
<tr>
<td>Reasoning</td>
<td>Arithmetic</td>
<td>8.74 (3.12)</td>
<td>10.49 (2.77)</td>
<td>11.00 (2.25)</td>
<td>4.78</td>
<td>.01</td>
<td>DlPFC &lt; non-dlPFC*</td>
</tr>
<tr>
<td>Reasoning</td>
<td>Matrix Reasoning</td>
<td>10.47 (3.22)</td>
<td>11.52 (2.85)</td>
<td>12.28 (2.94)</td>
<td>2.94</td>
<td>.06</td>
<td>None</td>
</tr>
</tbody>
</table>

Means are presented with SDs in parentheses. Significant between-group differences were determined with the Tukey’s HSD test. *$p < .05$; **$p < .01$.

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verbal items (Letter–Number Sequencing; \( p < .05 \), uncorrected). This pattern of findings suggests that the dlPFC may be critical for the manipulation of verbal knowledge in mathematical reasoning and working memory. To substantiate this conclusion, however, it is necessary to examine several factors that are relevant to the interpretation of the observed results: (1) anatomical specificity of the lesions, (2) lesion laterality, and (3) specificity of the cognitive deficit.

### 3.1. Focal dorsolateral prefrontal lesions

To strengthen the precision of our analysis, we examined the performance of a subset of patients in the dlPFC sample whose lesions were (1) confined to the frontal lobes, (2) lateralized, and (3) entailed damage within or adjacent to the peak area of dlPFC activation reported by a large-scale meta-analysis of fMRI studies on working memory (Wager and Smith, 2003). The results of this meta-analysis identified 86 peak activations reported by working memory studies within dlPFC, with a geometric center of activation in \( x = 40, y = 34, z = 29 \) (MNI coordinates; for further detail, see Wager and Smith, 2003). We assembled a left focal dlPFC sample \( (n = 7; \text{Fig. 2}) \) and a right focal dlPFC group \( (n = 9; \text{Fig. 3}) \) that each consisted of patients whose lesions were overlapping or adjacent to this peak activation site. In particular, five out of seven left dlPFC patients (Fig. 4a) and six out of nine right dlPFC patients (Fig. 4b) entailed damage to this region, supporting a more targeted assessment of the causal contribution of dlPFC to working memory. In addition, we further characterized the contribution of white matter pathways in the focal dlPFC samples, identifying that each patient group entailed damage within or adjacent to the (1) superior longitudinal fasciculus (primarily branch 1), (2) frontal aslant tract, and (3) fronto-striatal tracts (Thiebaut de Schotten et al., 2012; Mori et al., 2008).

### 3.2. Comparison with respect to prefrontal lesions

To further minimize differences between the dlPFC lesion patients and the brain-injured comparison group, we constructed a new comparison group consisting of patients with focal non-dlPFC lesions (focal non-dlPFC patient group; Supplementary Fig. 2; \( n = 20 \)). In contrast to the earlier non-dlPFC patient sample, this comparison group consisted of patients with lesions primarily confined to PFC rather than having lesions whose size and location were highly variable. As Supplementary Fig. 2 illustrates, the focal non-dlPFC sample entailed lesions primarily within ventromedial PFC, representing the subset of patients from the earlier non-dlPFC sample with highly focal lesions. Demographic and background cognitive function data for each patient group are presented in Table 3. No significant group differences were observed with respect to basic demographic variables (age, sex, years of education), pre- and post-combat measures of cognitive function, post-combat measures of verbal IQ and verbal comprehension, and lesion size. In summary, the focal patient groups were well matched with respect to (1) basic demographic variables, (2) pre- and post-combat measures of cognitive function, (3) lesion size, and (4) lesion location (i.e., primarily confined to PFC).

### 3.3. Specificity of cognitive deficit

To determine the effect of focal dlPFC lesions on components of working memory, we examined the performance of the left dlPFC \( (n = 7) \) and right dlPFC \( (n = 9) \) samples with respect to non-dlPFC lesion patients \( (n = 20) \) and neurologically healthy participants \( (n = 54) \). Because the assumptions underlying parametric statistics were not satisfied (e.g., homogeneity of variance, large sample size, and normality), non-parametric statistics were applied to test for group effects and for pairwise comparisons.

#### 3.3.1. Working memory

To summarize the results reported in Table 4 and Supplementary Figs. 3–22, focal lesions of the left or right dlPFC did not produce reliable deficits in working memory maintenance (Digit Span Forward and Spatial Span Forward), monitoring (Zero-Back), or under conditions of increasing cognitive load (One-, Two- and Three-Back). However,
Impairments in the left dlPFC patient group were observed for a test requiring the manipulation of verbal and auditory information in working memory (Letter–Number Sequencing) and approached significance for the test requiring the manipulation of non-verbal and spatial knowledge (Spatial Span Backward; \( p < .05 \), uncorrected). Additional analyses investigating the correlation between percent volume loss in dlPFC and performance on the administered tests of working memory revealed a converging pattern of findings (see Supplementary Table 2). In summary, our findings suggest that the left dlPFC is necessary for manipulating verbal/auditory and non-verbal/spatial information in working memory.

3.3.2. Reasoning
As Table 4 and Supplementary Figs. 3–24 illustrate, no reliable deficits were observed in the left dlPFC patient group for measures of mathematical (Arithmetic) or spatial reasoning (Matrix Reasoning). However, the right dlPFC patient group was significantly impaired for both neuropsychological tests of reasoning (for additional evidence, see Supplementary Table 2). This pattern of findings suggests that the right dlPFC is critical for manipulating information in the employed tests of arithmetic and spatial reasoning.

4. Discussion
The aim of the current investigation was to examine the necessity of the dlPFC for key elements of working memory. Using a relatively large sample of patients with dorsolateral prefrontal damage (\( n = 19 \)) and a wide-ranging assessment of cognitive function, we report several main findings. First, our results indicate that unilateral dlPFC is not necessary for working memory maintenance, monitoring, or for tasks that measure working memory performance under cognitive load. Second, our findings suggest that the dlPFC is important for manipulating representations in working memory (Letter–Number Sequencing) and in reasoning (Arithmetic; Table 2). Third, our results indicate that the left dlPFC is necessary for manipulating verbal and spatial knowledge in working memory (Letter–Number Sequencing; Spatial Span Backwards), while the right dlPFC is critical for the employed tests of verbal and spatial reasoning (Arithmetic; Matrix Reasoning; Table 4).

Our findings are therefore consistent with domain-general models of working memory, which posit that the dlPFC embodies specific computational mechanisms for monitoring and manipulating cognitive representations (Owen et al., 1996; Duncan and Owen, 2000; Miller and Cohen, 2001; Koechlin et al., 2003; Petrides, 2000, 2005).

A key contribution of our lesion study is to elucidate the nature of these mechanisms, demonstrating that the dlPFC is necessary for manipulating verbal and spatial knowledge. Functional neuroimaging evidence indicates that the dlPFC is selectively engaged in a wide range of working memory operations, with increased activation in this region observed (1) at the beginning of delayed-response trials in which the amount of to-be-remembered information approaches or exceeds short-term memory capacity, (2) during the subsequent delay interval when no information is accessible to the subject (Courtney et al., 1997; Zarahn et al., 1999), (3) for manipulating information during the delay period (D’Esposito and Postle, 1999; Postle et al., 1999; Rypma and D’Esposito, 1999), and (4) upon presentation of the probe stimulus, when a subject is required to select an appropriate response. These findings highlight the temporal dynamics of dlPFC function in working memory and suggest that this region is involved in several encoding- and response-related operations, as well as mnemonic and non-mnemonic processes that are engaged when manipulating information. The results of our lesion study demonstrate that although the dlPFC is associated with multiple cognitive operations, it is computationally necessary for the specific process of manipulating verbal and spatial knowledge.

The observed lateralization within the dlPFC further suggests that the left dlPFC supports manipulating representations in working memory and the right dlPFC supports the manipulation of information in a broader range of reasoning contexts. In both cases, the dlPFC implements specific
processes for manipulating cognitive representations (in the verbal and spatial domain) and therefore supports a domain-general model of the functional organization of the dLPFC. This pattern of findings is consistent with the proposal that the left dLPFC supports cognitive processes that are temporally bounded within working memory (Letter-Number Sequencing; Spatial Span Backwards), whereas the right dLPFC supports cognitive processes that extend beyond the scope of working memory and enable goal-directed behavior and adaptive decision making (Arithmetic; Matrix Reasoning; Barbey et al., 2009).

When evaluating the theoretical contributions of this study, it is important to emphasize the type of inferences that can be drawn from lesion data. While physiological studies of the nervous system are based on correlational methods (e.g., single- and multi-unit electrophysiology, electroencephalography, magnetoencephalography, measures of glucose metabolism and the blood oxygenation level-dependent response), lesion data support inferences about the necessity of a brain region for a given cognitive function. Interpretation of neuropsychological data, however, is subject to a different set of limitations. Lesion localization and the interruption of fibers of passage by a brain injury are often difficult to assess in human studies, and the damaged region may contribute in a non-specific way to the normal functioning of a distal region that is itself the true neural substrate of the function in question. It is important to emphasize that the dLPFC lesion patients under investigation here had damage within or adjacent to the (1) superior longitudinal fasciculus (primarily branch 1), (2) frontal aslant tract, and (3) fronto-striatal tracts (see Thiebaut de Schotten et al., 2012; Mori et al., 2008). Lesions within these white matter fiber tracts damage neural circuitry by disconnecting dLPFC and medial parietal cortex (superior longitudinal fasciculus), ventrolateral PFC and bilateral medial prefrontal (frontal aslant tract), and dLPFC and the dorsal striatum (fronto-striatal tracts). As a consequence, the observed pattern of working memory deficits reflects damage not only to the dLPFC but also to a crossroad of tracts that allow communication between several brain regions that have been implicated in working memory (for a meta-analytic review, see Owen et al., 2005).

Accumulating neuroscience evidence indicates that working memory and other higher cognitive processes centrally depend on white matter fiber tracts that synthesize information across a broadly distributed neural system. Recent voxel-based lesion-symptom mapping studies have sharpened our understanding of the role of white matter fiber tracts in binding the dLPFC and parietal cortex into an integrated system subserving working memory and general intelligence (Barbey et al., 2012a; Barbey et al., 2012b; Glascher et al., 2010, 2009; Chiang et al., 2009; Rudrauf et al., 2008). Barbey et al. (2012a) showed that the neural architecture of general intelligence and working memory is remarkably circumscribed, concentrated within the core of white matter fiber tracts that connect dLPFC with the inferior parietal cortex and that terminate in the superior parietal lobe. The observed reliance upon white matter fiber tracts suggests that working memory and other high-level cognitive processes are supported by the interregional communication among many brain areas, emphasizing the central role of the dLPFC and parietal cortex (Jung and Haier, 2007).

We emphasize, in closing, that understanding the neural architecture of working memory will ultimately require knowledge of the entire network of brain regions that participate, the contribution made by each component, and the role of white matter fiber tracts that communicate and synthesize information between them. The results of the present investigation contribute to this emerging research program by elucidating the involvement of the dLPFC, demonstrating that this region supports the manipulation of verbal and spatial representations in working memory. Although activation within the dLPFC is associated with a broad range of cognitive operations, our study indicates that this region is a central component of the neural systems underlying the manipulation of verbal and spatial knowledge.
Table 3 – Demographic and background data.

<table>
<thead>
<tr>
<th>Demographic data</th>
<th>L dlPFC</th>
<th>R dlPFC</th>
<th>Focal non-dlPFC</th>
<th>No lesion</th>
<th>Kruskal–Wallis $\chi^2$</th>
<th>Kruskal–Wallis $p$ value</th>
<th>Significant between-group differences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>57.57 (1.13)</td>
<td>60.00 (3.35)</td>
<td>58.52 (2.29)</td>
<td>59.52 (3.42)</td>
<td>7.42</td>
<td>.06</td>
<td>None</td>
</tr>
<tr>
<td>Sex (% male)</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>n/a</td>
<td>n/a</td>
<td>None</td>
</tr>
<tr>
<td>Years of education</td>
<td>15.43 (2.99)</td>
<td>14.22 (2.86)</td>
<td>14.33 (2.99)</td>
<td>15.19 (2.47)</td>
<td>5.75</td>
<td>.12</td>
<td>None</td>
</tr>
<tr>
<td>Pre-combat AFQT</td>
<td>53.43 (31.99)</td>
<td>54.88 (28.61)</td>
<td>54.20 (23.68)</td>
<td>65.40 (22.91)</td>
<td>7.06</td>
<td>.07</td>
<td>None</td>
</tr>
<tr>
<td>Post-combat AFQT</td>
<td>57.50 (29.49)</td>
<td>60.44 (25.89)</td>
<td>51.69 (22.68)</td>
<td>72.34 (22.99)</td>
<td>6.48</td>
<td>.09</td>
<td>None</td>
</tr>
<tr>
<td>Post-combat verbal IQ</td>
<td>107.57 (22.28)</td>
<td>101.11 (15.88)</td>
<td>102.70 (11.91)</td>
<td>109.66 (12.04)</td>
<td>3.70</td>
<td>.29</td>
<td>None</td>
</tr>
<tr>
<td>Total percent volume loss (cm$^3$)</td>
<td>2.99 (1.42)</td>
<td>2.55 (2.03)</td>
<td>2.97 (1.90)</td>
<td>n/a</td>
<td>1.19</td>
<td>.55</td>
<td>None</td>
</tr>
</tbody>
</table>

Data are presented as means with SDs in parentheses. “Age” refers to age at the time of Phase 3 evaluation. “Sex” refers to the percent of male veterans. “Years of education” refers to the total number of years of education the veterans completed. “Pre-combat AFQT” refers to index scores on the Armed Forces Qualification Test, a battery of tests measuring basic cognitive function at the time of enlistment (pre-injury). “Post-combat AFQT” refers to index scores on the Armed Forces Qualification Test administered at Walter Reed Medical Center after injury. “Post-combat verbal IQ” refers to the Phase 3 verbal IQ index score from the WAIS. There were no significant differences among groups for any measure. Non-parametric statistics were used to test for group effects and for the pairwise comparisons given the small number of participants in each sample. Significant between-group differences were determined with the Mann–Whitney U test.

Table 4 – Neuropsychological tests of working memory and reasoning.

<table>
<thead>
<tr>
<th>Cognitive function</th>
<th>Cognitive measure</th>
<th>L dlPFC</th>
<th>R dlPFC</th>
<th>Focal non-dlPFC</th>
<th>No lesion</th>
<th>Kruskal–Wallis $\chi^2$</th>
<th>Kruskal–Wallis $p$ value</th>
<th>Significant between-group differences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maintenance</td>
<td>Digit Span Forward</td>
<td>6.00 (1.41)</td>
<td>5.89 (1.05)</td>
<td>6.40 (1.19)</td>
<td>6.68 (1.22)</td>
<td>4.83</td>
<td>.19</td>
<td>None</td>
</tr>
<tr>
<td>Maintenance</td>
<td>Spatial Span Forward</td>
<td>8.57 (2.94)</td>
<td>9.11 (3.89)</td>
<td>9.89 (2.56)</td>
<td>10.22 (2.7)</td>
<td>3.05</td>
<td>.39</td>
<td>None</td>
</tr>
<tr>
<td>Monitoring: no cognitive load</td>
<td>Zero-Back Errors</td>
<td>1.00 (.71)</td>
<td>2.56 (3.47)</td>
<td>1.16 (1.89)</td>
<td>1.08 (2.33)</td>
<td>3.09</td>
<td>.38</td>
<td>None</td>
</tr>
<tr>
<td>Monitoring: low cognitive load</td>
<td>One-Back Errors</td>
<td>3.40 (2.97)</td>
<td>3.67 (3.24)</td>
<td>3.00 (2.75)</td>
<td>2.57 (2.48)</td>
<td>1.26</td>
<td>.74</td>
<td>None</td>
</tr>
<tr>
<td>Monitoring: medium cognitive load</td>
<td>Two-Back Errors</td>
<td>5.20 (1.92)</td>
<td>4.89 (2.37)</td>
<td>5.26 (2.81)</td>
<td>3.85 (2.39)</td>
<td>6.66</td>
<td>.08</td>
<td>Focal non-dlPFC &gt; no lesion**</td>
</tr>
<tr>
<td>Monitoring: high cognitive load</td>
<td>Three-Back Errors</td>
<td>6.00 (2.74)</td>
<td>5.67 (3.61)</td>
<td>6.42 (2.65)</td>
<td>4.68 (2.40)</td>
<td>5.27</td>
<td>.15</td>
<td>Focal non-dlPFC &gt; no lesion**</td>
</tr>
<tr>
<td>Manipulation</td>
<td>Letter–Number Sequencing</td>
<td>8.14 (2.91)</td>
<td>9.44 (3.68)</td>
<td>9.95 (2.95)</td>
<td>11.04 (2.66)</td>
<td>7.73</td>
<td>.05</td>
<td>L dlPFC &lt; no lesion**</td>
</tr>
<tr>
<td>Manipulation</td>
<td>Digit Span Backward</td>
<td>4.14 (1.22)</td>
<td>4.67 (1.41)</td>
<td>4.75 (1.21)</td>
<td>4.96 (1.41)</td>
<td>2.49</td>
<td>.48</td>
<td>None</td>
</tr>
<tr>
<td>Manipulation</td>
<td>Spatial Span Backward</td>
<td>8.86 (3.81)</td>
<td>11.33 (2.96)</td>
<td>11.28 (2.49)</td>
<td>12.02 (3.12)</td>
<td>6.02</td>
<td>.11</td>
<td>None</td>
</tr>
<tr>
<td>Reasoning</td>
<td>Arithmetic</td>
<td>8.71 (3.82)</td>
<td>8.22 (3.11)</td>
<td>10.40 (2.30)</td>
<td>11.00 (2.25)</td>
<td>8.70</td>
<td>.03</td>
<td>R dlPFC &lt; no lesion**</td>
</tr>
<tr>
<td>Reasoning</td>
<td>Matrix Reasoning</td>
<td>10.57 (3.78)</td>
<td>9.44 (2.79)</td>
<td>11.89 (2.47)</td>
<td>12.28 (2.94)</td>
<td>7.94</td>
<td>.04</td>
<td>R dlPFC &lt; focal non-dlPFC**</td>
</tr>
</tbody>
</table>

Means are presented with SDs in parentheses. Because the assumptions of parametric statistics were not satisfied (Supplementary Table 1, Supplementary Figs. 1–22), non-parametric statistics were used to test for group effects and for the pairwise comparisons. Significant between-group differences were determined with the Mann–Whitney U test. Neuroscience evidence supporting the involvement of the dlPFC in working memory motivated the assessment of between-group differences when the Kruskal–Wallis test did not reach significance (see Introduction).

*p < .05; **p < .01.

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Supplementary material

Supplementary material associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.cortex.2012.05.022.

REFERENCES


