Prefrontal Cortical Representation of Visuospatial Working Memory in Monkeys Examined by Local Inactivation With Muscimol

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Sawaguchi, Toshiyuki and Michio Iba. Prefrontal cortical representation of visuospatial working memory in monkeys examined by local inactivation with muscimol. J Neurophysiol 86: 2041–2053, 2001. In primates, dorsolateral areas of the prefrontal cortex (PFC) play a major role in visuospatial working memory. To examine the functional organization of the PFC for representing visuospatial working memory, we produced reversible local inactivation, with the local injection of muscimol (5 μg, 1 μl), at various sites (n = 100) in the dorsolateral PFC of monkeys and observed the behavioral consequences in an oculomotor delayed-response task that required memory-guided saccades for locations throughout both visual fields. At 82 sites, the local injection of muscimol induced deficits in memory-guided saccades to a few specific, usually contralateral, target locations that varied with the location of the injection site. Such deficits depended on the delay length, and longer delays were associated with larger deficits in memory-guided saccades. The injection sites and affected spatial locations of the target showed a gross topographical relationship. No deficits appeared for a control task in which the normal function of a small region of frontal cortical areas of monkeys, such as the premotor cortex (Kurata and Hoffman 1994) and the frontal eye field (Dias and Segraves 1999; Sommer and Tehovnik 1997). We obtained evidence that memoranda for specific visuospatial coordinates are represented in a “memory map” within the dorsolateral PFC to represent visuospatial working-memory processes.

METHODS

Subjects and behavioral tasks

Two male rhesus monkeys (Macaca mulatta, 6–7 kg) were used in the present study. Throughout the experiment, the subjects were treated in accordance with the Guide for Care and Use of Laboratory Animals (National Institutes of Health) and the Guide for Care and Use of Laboratory Animals of our institute. The monkeys were trained to perform an ODR task and a control (CON) task. Before training, each monkey was habituated to a monkey chair, and then preliminary surgery was performed under deep anesthesia with pentobarbital sodium (~25 mg/kg iv) and aseptic conditions. Two hollow rods (8 mm ID), for restraining the head of the monkey, were implanted on the anterior and posterior portions of the skull with dental acrylic. Smaller stainless-steel bolts (3 mm ID), for grounding, were anchored to the marginal portions of the skull and fixated with dental acrylic. To prevent infection, antibiotics were injected intramuscularly on the day of the surgery and daily for 1 wk afterward. A few weeks after surgery, each monkey began training on the ODR and CON tasks. During daily sessions, the monkey sat in a monkey chair and faced a multiscan 21-in cathode ray tube (CRT) monitor (PC-TV471, NEC, Tokyo) placed ~60 cm in front of him. The monkey’s head was rigidly fixated by two stainless steel bars (8 mm diam) to a stereotaxic frame located at the top of the monkey chair, and a water spout was positioned close to the monkey’s mouth. The task and the recordings were controlled by a system consisting of an infrared eye-camera system (R-21C-A, RMS, Hiroasaki, Japan), two personal computers (PC9801FE and PC9801BX, NEC, Tokyo) that were networked by RS232C and parallel I/O, and other associated peripherals. The eye-camera system was

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connected to the personal computers via A/D converters (AB98-05A/4, ADTEK, Yokohama, Japan) and was used for monitoring and sampling eye positions. One of the personal computers (PC9801FE) controlled the task and generated visual stimuli on the CRT monitor, and the other computer (PC9801BX) monitored and sampled eye positions and task events. The computer programs were written in C and partially in Assembly languages.

The phases of the ODR and CON tasks are illustrated in Fig. 1A. The ODR task was started when the monkey fixated on a central spot (a white square, 0.2 × 0.2°) on the CRT. One second later, a visual cue (white square, 0.5 × 0.5°) was presented for 0.5 s, and this was followed by a delay period. The cue was presented randomly at 1 of 16 peripheral locations; their directions were from 0 (right) to 360° counter-clockwise at 45° intervals, with eccentricities of 10 and 20° (Fig. 1B). After a delay period of 2–6 s (usually 4 s), the fixation spot was then extinguished (“go” signal), which instructed the monkey to make a memory-guided saccade to the location that had been cued prior to the delay period. Correct responses were defined as eye movement which fell within a diameter of 5–10° around the target location. A correct response was rewarded by a drop of water 0.2 s after the response. When the eye movements did not fall within the prescribed window or when the monkey failed to respond within the allotted 0.7 s, the response was defined as an error and was not rewarded. Trials were separated by an intertrial interval of 2 s. The CON task was exactly the same as the ODR task except that the visual cue remained on during the “delay” and response periods, and the subject made a visually guided saccade to the visible target. The monkeys performed ~500–700 trials for each task during daily training or recording sessions. The correct-response rate was 95% in the final stages of training sessions and in the predrug period of daily recording sessions. In the predelay period of a typical recording session, the latency of saccades from the onset of the go signal was ~180–220 ms with SDs of ~10–25 ms for the ODR task and ~200–240 ms with SDs of ~15–30 ms for the CON task; this onset latency varied somewhat by the target location and by the subject.

Experimental procedures and data analysis

After the training was completed, surgery for the experiment was performed under aseptic conditions as in the preliminary surgery.

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**FIG. 1.** The behavioral task and injection sites. A: the oculomotor delayed-response (ODR) and control (CON) tasks. In the ODR task, monkeys fixated on a fixation spot on a CRT monitor, and a visual cue came on for 0.5 s, followed by a delay period. After the delay period (usually 4 s), the fixation spot turned off, instructing the monkeys to make a memory-guided saccade (arrow) to the location that had been indicated by the cue prior to the delay. The correct response was rewarded by a drop of water 0.2 s after the response. Trials were separated by an intertrial interval (ITI) of 2 s. The CON task was exactly the same as the ODR task except that the cue remained on during the “delay” period, and the monkeys made a visually guided saccade to the visible target at the end of the “delay” period. B: locations of cue presentation. The central square indicates the fixation spot. The cue was presented at 16 locations, which were separated by 45° with eccentricities of 10 and 20°. C: tracing of injection sites on the surface of the dorsolateral prefrontal cortex (PFC) of the 2 monkeys (SA and TA). Muscimol (MUS) was injected at a total of 100 sites in the dorsolateral PFC and at 8 sites in the frontal eye field (FEF). The injection sites associated with deficits in the ODR task by MUS are indicated (●), - ineffective sites; ▲, injection sites in the FEF. PS, principal sulcus; AS, arcuate sulcus.
Each monkey was anesthetized with pentobarbital sodium (~25 mg/kg iv), and an oval opening was made in the skull to expose the dura over the frontal cortex. An oval cylinder (18 × 36 mm ID) was positioned over the opening and fitted in place with dental acrylic. Prophylactic antibiotics were injected intramuscularly on the day of surgery and daily for 1 wk afterward.

A few weeks after surgery, the experimental session was started. While the monkeys performed the ODR and CON tasks, 1 μl of muscimol hydrochloride (MUS, 5 μg/μl, dissolved in 0.9% saline) was locally injected into the dorsolateral PFC using a microsyringe (MS-NP10; Ito, Osaka, Japan). The methods of injection were similar to those in our previous study for the monkey frontal cortex (Sawaguchi et al. 1996). Briefly, injections were made 3–5 mm from the dural surface. The needle (0.52 mm OD, 0.13 mm ID, ~0.008 mm² open surface area) had a sharp angle (~27°) at the tip, and the opening of the tube was located ~0.5 mm from the tip. The needle was positioned using a micromanipulator (MO-95, Narishige, Tokyo) and a plastic grid with numerous small holes (0.7 mm ID, 1.5 mm apart from each other) attached to the cylinder. This method allowed us to precisely control the location of the insertion with an accuracy of ~0.2 mm and its relocation in subsequent sessions. Furthermore, the needle of the syringe was coated with polyurethane, and multiple neuronal activities at the injection site were recorded using the tip (resistance, 0.2–0.5 MΩ). Thus we were able to confirm that the tip of the needle was located within the gray matter of the cortex.

The experimental sessions consisted of blocks of trials 5 min long. In each block, the monkey performed the ODR or CON task, and blocks associated with the CON task were usually intermixed with those associated with the ODR task. The monkeys performed four or six blocks for each task (i.e., 40 or 60 min) before the injection of MUS. After the predrug period, testing was interrupted for 2–3 min while 1 μl of MUS was injected at a rate of ~0.5 μl/min, and postdrug trials were started. The postdrug period consisted of at least eight blocks of each task (i.e., 80 min) after the injection. The following behavioral parameters were examined and compared between the pre- and postdrug trials: discrepancy between the target location and the end point of the saccade during the response period; the onset latency of the response after the onset of the go signal; two-dimensional trajectories of saccades; and the amplitude and latency of saccades. The predrug blocks were combined into one score for each task, and this value was compared with every postdrug block by one-way ANOVA followed by the Newman-Keuls procedure for multiple comparisons. Furthermore, since MUS usually produced a significant effect within a few minutes after injection, which lasted for ~40–60 min after injection, and the predrug control level was gradually restored during the experimental sessions, although this level was not quite achieved. Furthermore, the effects of MUS on ODR performance were sensitive to the length of the delay period: longer delays were associated with larger deficits in ODR performance, although this level was not quite achieved. Furthermore, the effects of MUS on ODR performance were sensitive to the length of the delay period: longer delays were associated with larger deficits in ODR performance, although this was examined in only a few sessions (n = 8).

**Effects of local injections of MUS on ODR and CON performance**

The injection of MUS affected task performance for 82 of the 100 injection sites in the dorsolateral PFC. The results for one such site are shown in Fig. 2. In this case, MUS (5 μg, 1 μl) was injected into the left dorsolateral PFC (Fig. 2, inset). Prior to injection, the monkey performed the ODR and CON tasks for ~60 min without error. The length of the delay period was 4 s, and the visuospatial cue was randomly presented at 1
of 16 locations (8 directions and 2 eccentricities of 10 and 20°). Figure 2A shows percent changes, compared with the predrug period, in mean values of the discrepancy during the postdrug period of 60 min after injection for each target location. Data are from 60 min before injection and from 0 to 60 min after injection. Squares with dashed lines indicate 0% changes, and larger squares indicate greater percentage increases. The shaded square is the target location for which the discrepancy increased significantly after injection \( (P < 0.01, \text{Mann-Whitney’s } U \text{ test}) \). The deficit was limited to a specific target location (10° in eccentricity, 0° in direction) during ODR task performance. No deficit appeared in performance of the CON task, which required only visually guided saccades. B: horizontal (H) and vertical (V) traces of eye movements from the onset of the go signal, and 2-dimensional trajectories (T) of saccades to the 0° target location (10° in eccentricity) for which memory-guided saccades were significantly affected, before (Pre) and after (Post) injection.

To examine the time course of the effect induced by MUS injection, the accuracy of saccades before and after injection was calculated for the injection described in the preceding text. Figure 3A shows the results in which the discrepancy for a specific target location (0° direction and 10° eccentricity) is illustrated. As shown, the discrepancy in the memory-guided saccade increased within 20 min after MUS injection, reached a peak at \( \sim 40 \text{ min} \), and gradually returned to the predrug control level, although the discrepancy did not reach the control level within this session.

To examine whether or not the velocity of the memory-guided saccade was affected by MUS, the relationship between the amplitude and the peak velocity of the saccades was examined. In Fig. 3B, velocity is plotted against amplitude for memory-guided saccades in ODR trials before (○) and within 60 min after MUS injection (●). Only the memory-guided saccades to the two targets located in the 0° direction (eccentricities, 10 and 20°) were examined because this direction was the most strongly affected by
injection. As shown in Fig. 3B, the velocity of the memory-guided saccades did not change after injection.

In contrast to the ODR task, performance in the CON task was not affected by MUS injection: performance in the CON task was 100% correct for all target locations throughout the experimental session, and the discrepancy did not significantly change for any target location after injection (Fig. 2A), including the target located at 0° (10° eccentricity), for which memory-guided saccades were impaired (1.36 ± 0.65°, n = 7, for the predrug trial; 1.25 ± 0.39°, n = 11, for the postdrug trial; P > 0.05, NS, U test). The trajectories of visually guided saccades to this target location were similar before and after injection (Fig. 2B). Furthermore, the accuracy of the visually guided saccade in the CON task was constant throughout the experimental period (Fig. 3A).

Another example of the effect of MUS injection into the dorsolateral PFC on task performance is illustrated in Fig. 4. MUS was injected into a site close to the rostral edge of the upper arcuate sulcus, and errors in ODR performance appeared for a specific target in the upper-right (45°) visual field with 20° eccentricity (Fig. 4A). Memory-guided saccades toward this target location were misdirected or distorted after injection.

**FIG. 3.** A: time courses of the change in the discrepancy between the target location and the end point of the saccade for the ODR and CON tasks. Data are the same as those in Fig. 2, and data for the target location (inset) that was significantly affected by MUS injection are shown. Means and SE (bar) based on 3–7 trials per time point. B: relationships between amplitude and peak velocity of memory-guided saccades in the ODR task before and after MUS injection. The data are the same as those for Fig. 2, and the data for target locations in the 0° direction (eccentricities of 10 and 20°) (inset) are shown. Each point represents a memory-guided saccade. ○, data before injection; ●, data within 60 min after the injection. The relationship between the amplitude and velocity of the saccade was similar before and after the injection.

**FIG. 4.** Effects of MUS injection on performance in the ODR and CON tasks. MUS (5 μg/μl, 1 μl) was injected into a site in the left PFC (inset) while the monkey (SA) performed the ODR and CON tasks. Format as in Fig. 2.
The discrepancy for this target location significantly increased after injection (3.84 ± 1.07°, n = 11, for the predrug trial; 11.08 ± 6.23°, n = 15, for the postdrug trial; P < 0.01, U test). In contrast, performance in the CON task was not affected by MUS. The discrepancy did not significantly change for any target location after injection (Fig. 4A), including the target location (45° in direction and 20° in eccentricity) for which memory-guided saccades were impaired (2.92 ± 0.84°, n = 10, for the predrug trial; 3.17 ± 1.02°, n = 13, for the postdrug trial; P > 0.05, NS, U test). The trajectories of visually guided saccades to this target location were similar before and after injection (Fig. 4B).

Thus local injections into the dorsolateral PFC induced deficits in ODR performance unaccompanied by deficits in the CON task. In the ODR task, affected targets were limited to a specific location, and the trajectories of the memory-guided saccades to such targets were abnormal after injection; they were frequently misdirected or distorted.

**Delay dependency**

The effect of MUS depended on the delay length: longer delays were associated with larger deficits in ODR performance for all eight injection sites examined. An example is shown in Fig. 5. In this case, MUS was injected into a site in the left dorsolateral PFC of monkey TA while it was performing the ODR task with delay periods of 2, 4, and 6 s. As shown in Fig. 5, the largest deficit appeared at the 315° target location with 20° eccentricity for trials with each delay. The trajectories of memory-guided saccades toward this target location were more strongly impaired for trials with longer delays. The percent increase in the discrepancy for the affected target was the smallest for trials with a 2-s delay, followed by trials with a 4-s delay; the largest increase in discrepancy was observed with a 6-s delay. The mean discrepancy for the affected target location during predrug trials was 3.34 ± 1.36° (n = 8) for a 2-s delay, 3.64 ± 1.43° (n = 8) for a 4-s delay, and 2.81 ± 1.41° (n = 7) for a 6-s delay, with no significant differences among trials with different delays (ANOVA, df = 2, 20, F = 0.666, P = 0.525, NS). The mean discrepancy during postdrug trials was 10.13 ± 8.04° (n = 8) for a 2-s delay, 17.37 ± 11.01° (n = 8) for a 4-s delay, and 24.02 ± 10.27° (n = 12) for a 6-s delay, with a significant difference among trials with different delays (ANOVA, df = 2, 27, F = 6.054, P < 0.01). The discrepancy for a 2-s delay was significantly smaller than those for a 4-s delay (P < 0.05, U test) and a 6-s delay (P < 0.01), although there was no significant difference between trials with 4- and 6-s delays (P > 0.05, NS).

**Dose dependency**

The effect of MUS was dose dependent: a larger dose of MUS (10 μg, 2 μl) induced larger deficits in ODR performance, compared with the standard dose (5 μg, 1 μl), for all five injection sites examined; no deficits occurred for CON performance even with the highest dose of MUS in all of the injection sites. For the case shown in Fig. 6, the largest deficit in the ODR task appeared for the right (0°) target direction with 10° eccentricity when the standard dose (5 μg, 1 μl) of MUS was injected (Fig. 6A). For this target location, the discrepancy was significantly increased...
(2.67 ± 1.72°, n = 14, for the predrug trial; 7.76 ± 5.99°, n = 10, for postdrug trials; P < 0.01, U test), and memory-guided saccades were misdirected or distorted (Fig. 6A). No other target location was affected by injection, as with other cases examined with the standard dose of MUS. When a higher dose (10 μg, 2 μl) of MUS was injected in a different daily session (Fig. 6B), memory-guided saccades for the same target location (0° in direction, 10° in eccentricity) were impaired: the discrepancy for this target location was significantly increased after injection (1.79 ± 1.24°, n = 8, for the predrug trial; 7.36 ± 5.80°, n = 12, for postdrug trials; P < 0.01, U test). Furthermore, memory-guided saccades were also impaired for three target locations surrounding this target location; i.e., the upper-right (45°) location with 10° eccentricity, right (0°) location with 20° eccentricity, and lower-right (315°) location with 10° eccentricity. The discrepancy values (predrug vs. postdrug trials) were 2.13 ± 0.99° (n = 10) versus 7.63 ± 4.54° (n = 12; P < 0.01) for the 45° direction with 10° eccentricity, 3.55 ± 1.46° (n = 8) versus 9.74 ± 6.43° (n = 11; P < 0.05) for the 0° direction with 20° eccentricity, and 1.93 ± 1.47° (n = 7) versus 6.59 ± 5.44° (n = 13; P < 0.05) for the 315° direction with 10° eccentricity. Thus a higher dose of MUS induced deficits in memory-guided saccades for additional target locations surrounding the particular target location that was impaired with the standard dose of MUS.

Relationships between injection sites and target locations affected

There was a constant relationship between injection sites and target locations that were affected in the ODR task by MUS injection, as shown in the example in Fig. 7. In this figure, the percent change in discrepancy in memory-guided saccades in
ODR performance is illustrated for four typical injection sites in the left hemisphere of monkey SA. Injection into a caudal site close to the principal sulcus affected the memory-guided saccade toward the upper, peripheral target location (90° direction, 20° eccentricity); injection into a site close to the middle of the principal sulcus affected the memory-guided saccade toward the lower-right, peripheral target location (315° direction, 20° eccentricity); injection into a more rostral site in the dorsal convexity affected the memory-guided saccade toward the lower, more central target location (270° direction, 10° eccentricity); and injection into a caudal site of the dorsolateral PFC affected the memory-guided saccade toward the upper-right, more central target location (45° direction, 10° eccentricity).

Figure 8 summarizes the relationships between each injection site and the direction and eccentricity of the target location that was the most strongly affected by each injection for three hemispheres of the two monkeys (SA and TA). In all three hemispheres, the deficits in memory-guided saccades caused by MUS injection were limited to a few specific, usually contralateral, target locations. In the left hemisphere of monkey SA, injection into rostral sites of the dorsolateral PFC induced deficits in memory-guided saccades toward lower target locations, whereas injection into caudal sites affected memory-guided saccades to upper locations. Injection into lateral sites affected memory-guided saccades to a peripheral target location (i.e., 20° in eccentricity), and injection into more medial sites affected memory-guided saccades to a more central location (10° in eccentricity). The injection sites associated with deficits for horizontal target locations (i.e., 0°) were arranged from medial to lateral, and those associated with deficits in vertical target locations (90 or 270°) appeared to be arranged from caudal to rostral. Similar topographic relationships were found in both hemispheres of the other monkey TA (TA-L and TA-R in Fig. 8).

Injections into the FEF

As a comparison, MUS was injected into a total of eight sites in the FEF, although we did not extensively test in the FEF. In all of the injection sites, MUS affected saccades in both ODR and CON tasks, and a typical example is shown in Fig. 9. In this case, MUS was injected into a site close to the arcuate sulcus of the left hemisphere, where ICMS induced eye movement. MUS injection affected memory-guided saccades during ODR performance as well as visually guided saccades during CON performance; the effect was the most significant for lower-right target locations (225°), in particular, with 20° eccentricity (Fig. 9A). Memory-guided saccades toward this target location (i.e., 225°, 20° eccentricity) were misdirected or distorted after injection, as were visually guided saccades, but not to the same extent; smaller saccades were more frequently observed after injection (Fig. 9B).

Indeed, the amplitude of visually guided saccades toward this location in the CON task was significantly shorter for postdrug trials (17.56 ± 2.62°, n = 9) than for predrug trials (20.92 ± 1.26°, n = 11; P < 0.01, U test). The discrepancy for this target location significantly increased after injection for both the ODR and CON tasks (2.16 ± 1.15°, n = 7, for the predrug trial; 8.96 ± 6.72°, n = 10, for the postdrug trial; P < 0.01, for ODR; and
1.91 ± 1.15°, n = 9, for the predrug trial; 4.04 ± 1.88°, n = 11, for the postdrug trial; P < 0.05, for CON). The onset latency also significantly increased after injection for the both the ODR and CON tasks (257 ± 19 ms for the predrug trial; 345 ± 73 ms for the postdrug trial; 265 ± 27 ms for the predrug trial; 380 ± 53 ms for the postdrug trial; P < 0.01, for CON). Further, the velocities of both memory- and visually guided saccades appeared to become slower after injection, as evident in Fig. 10, in which the peak velocity is plotted against amplitude for saccades in the ODR (left) and CON (right) trials before (○) and after MUS injection (●). Thus MUS injection into the FEF increased the discrepancy and onset latency of both memory- and visually guided saccades and made both saccades slower.

DISCUSSION

Involvement of a specific site in the PFC in working memory for a specific visuospatial coordinate

In the present study, we locally injected MUS (usually 5 μg, 1 μl) at various sites in the dorsolateral PFC of monkeys and observed the behavioral consequences in an ODR task. The ODR task required memory-guided saccades for locations throughout both visual fields and, hence, required working memory for visuospatial information (Funahashi et al. 1989). Since the intracerebral injection of a few microliters of MUS suppresses the activity of neurons within a few cubed millimeters of the injection site for some hours (cf. Kurata and Hoffman 1994), the present method is useful for mapping the normal function of a small region of the cerebral cortex. One microliter of solution injected into cerebral tissue has been demonstrated to spread ~1 mm in diameter (Myers 1966).

The local injection of 1 μl MUS induced deficits in memory-guided saccades to a few specific, usually contralateral, target locations that varied with the location of the injection site. The deficit was characterized by misdirection/distortion of memory-guided saccades and a decrease in accuracy as measured by the discrepancy between the end point of the saccade relative to the remembered target location. No deficits appeared for the CON task in which the subject was required to make a visually guided saccade to a visible target. The CON task required sustained attention, and all of the perceptual and motor features

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of the ODR task, but did not require working memory. Therefore factors other than the mnemonic demand, such as deficits in global attention, sensory perception, or execution of movements, can be excluded as explanations for the deficit in ODR performance, suggesting that such deficits specifically involve the working memory process. Furthermore, the effect of MUS depended on the delay length, and longer delays were associated with larger deficits in ODR performance. In contrast to the

**FIG. 9.** Effects of MUS injection into the frontal eye field (FEF) on performance in the ODR and CON tasks. MUS (5 μg/μl, 1 μl) was injected into a site in the left FEF (inset) while the monkey (SA) performed the ODR and CON tasks. Format as in Fig. 2. The eye traces (B) are for saccades to the 225° target location (20° in eccentricity) for which saccades were the most affected by the injection.

**FIG. 10.** Relationships between amplitude and peak velocity of memory-guided saccades in the ODR task and visually guided saccades in the CON task before and after MUS injection into the FEF. The data are the same as those for Fig. 9, and the data for target locations in the 225° direction (eccentricities of 10° and 20°; inset) are shown. ○, data before injection; ●, data within 60 min after injection.
dorsolateral PFC, MUS injections into the FEF affected both memory-guided saccades in the ODR task and visually guided saccades in the CON task, which is consistent with a previous study with MUS injection into the FEF (Dias and Segraves 1999); the FEF is considered to be involved in the voluntary saccade-generation/control mechanism itself, as has been emphasized by other authors (e.g., Bruce et al. 1985; Dias and Segraves 1999). These findings emphasize the basic mnemonic function of the dorsolateral PFC: it is essential for guiding behavior by internal cues, rather than by external sensory cues. We previously dissociated mnemonic and sensorimotor components of the ODR in our study of D1 dopamine receptor antagonism in the dorsolateral PFC (Sawaguchi and Goldman-Rakic 1991, 1994), which is consistent with the present finding. Thus a specific site in the dorsolateral PFC appears to be responsible for the working memory process for a specific visuospatial coordinate to guide goal-directed behavior.

“Memory map” representation in the PFC

One of the striking findings in the present study is that the injection sites and affected spatial locations of the target showed a gross topographic relationship; injections into more caudal sites of the dorsolateral PFC induced deficits in memory-guided saccades to upper target locations, and injections into more lateral sites affected memory-guided saccades to more peripheral target locations. The affected target locations were mainly located in the visual field contralateral to the injection site. These findings suggest that the dorsolateral PFC of each hemisphere contains a topographical memory map for representing visuospatial working memory, mainly for the contralateral visual field, as schematically illustrated in Fig. 11. The memory map in each hemisphere appears to contain a horizontal meridian of the memory field oriented perpendicular to the principal sulcus, and a vertical meridian on the dorsomedial convexity that might form a U-shaped curve surrounding the horizontal meridian. This conclusion is consistent with evidence from studies using physiological recordings (Funahashi et al. 1989), surgical lesions (Funahashi et al. 1993), and chemical dysfunction with dopamine antagonists (Sawaguchi and Goldman-Rakic 1990, 1994) that individual neurons of the dorsolateral PFC have memory fields, mainly for the contralateral visual field, and appear to be distributed topographically, although neuronal data on this point are still scarce.

In previous studies using bicuculline methiodide (BMI), a GABA antagonist, and a manual delayed-response (DR) task with left-right cues in monkeys, injections of BMI into the dorsolateral PFC induced directional errors in the DR task (Sawaguchi et al. 1988, 1989). The effective site for BMI injection was limited to the bottom of the principal sulcus. Based on the results of the present study, the memory map appears to be represented mainly in the dorsal convexity of the PFC, in which BMI injections did not induce any directional errors in previous studies. The discrepancy between the present and previous studies may be due to the fact that the previous studies did not control eye movements of monkeys, and the subjects were not required to maintain fixation during trials. Under these conditions, the monkey may be able to receive visuospatial cues in a given visual field and no deficits occurred following BMI injection into any sites in the dorsal convexity that are associated with specific visuospatial coordinates in the memory map. It is also possible, and more likely, that injection sites in the dorsolateral convexity in the previous studies were limited and outside of sites that are specifically associated with the left-right coordinate (with eccentricities of 10 and 20°) in the memory map. In addition, there is another possibility that the memory map in the dorsal convexity is just for eye movement, although this is less likely since the dorsal convexity has connections with nonprimary motor cortical areas, such as the premotor cortex (Barbas et al. 1999; Watanabe-Sawaguchi et al. 1991). On the other hand, the present study did not explore the depth of the principal sulcus, and it is unknown whether or not there is an additional memory map in the principal sulcal cortex. However, previous lesion studies have demonstrated that the principal sulcal area is critical for manual DR (Gross and Weiskrantz 1962) and ODR (Funahashi et al. 1993) tasks, and the BMI-injection studies indicated that the bottom of the principal sulcus is critical for the manual DR task. Therefore it is plausible that an additional memory map might be represented in the principal sulcal cortex. This problem should be examined by further studies.

“Memory map” representation in the PFC

One of the striking findings in the present study is that the injection sites and affected spatial locations of the target showed a gross topographic relationship; injections into more caudal sites of the dorsolateral PFC induced deficits in memory-guided saccades to upper target locations, and injections into more lateral sites affected memory-guided saccades to more peripheral target locations. The affected target locations were mainly located in the visual field contralateral to the injection site. These findings suggest that the dorsolateral PFC of each hemisphere contains a topographical memory map for representing visuospatial working memory, mainly for the contralateral visual field, as schematically illustrated in Fig. 11. The memory map in each hemisphere appears to contain a horizontal meridian of the memory field oriented perpendicular to the principal sulcus, and a vertical meridian on the dorsomedial convexity that might form a U-shaped curve surrounding the horizontal meridian. This conclusion is consistent with evidence from studies using physiological recordings (Funahashi et al. 1989), surgical lesions (Funahashi et al. 1993), and chemical dysfunction with dopamine antagonists (Sawaguchi and Goldman-Rakic 1990, 1994) that individual neurons of the dorsolateral PFC have memory fields, mainly for the contralateral visual field, and appear to be distributed topographically, although neuronal data on this point are still scarce.

In previous studies using bicuculline methiodide (BMI), a GABA antagonist, and a manual delayed-response (DR) task with left-right cues in monkeys, injections of BMI into the dorsolateral PFC induced directional errors in the DR task (Sawaguchi et al. 1988, 1989). The effective site for BMI injection was limited to the bottom of the principal sulcus. Based on the results of the present study, the memory map appears to be represented mainly in the dorsal convexity of the PFC, in which BMI injections did not induce any directional errors in previous studies. The discrepancy between the present and previous studies may be due to the fact that the previous studies did not control eye movements of monkeys, and the subjects were not required to maintain fixation during trials. Under these conditions, the monkey may be able to receive visuospatial cues in a given visual field and no deficits occurred following BMI injection into any sites in the dorsal convexity that are associated with specific visuospatial coordinates in the memory map. It is also possible, and more likely, that injection sites in the dorsolateral convexity in the previous studies were limited and outside of sites that are specifically associated with the left-right coordinate (with eccentricities of 10 and 20°) in the memory map. In addition, there is another possibility that the memory map in the dorsal convexity is just for eye movement, although this is less likely since the dorsal convexity has connections with nonprimary motor cortical areas, such as the premotor cortex (Barbas et al. 1999; Watanabe-Sawaguchi et al. 1991). On the other hand, the present study did not explore the depth of the principal sulcus, and it is unknown whether or not there is an additional memory map in the principal sulcal cortex. However, previous lesion studies have demonstrated that the principal sulcal area is critical for manual DR (Gross and Weiskrantz 1962) and ODR (Funahashi et al. 1993) tasks, and the BMI-injection studies indicated that the bottom of the principal sulcus is critical for the manual DR task. Therefore it is plausible that an additional memory map might be represented in the principal sulcal cortex. This problem should be examined by further studies.

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relations to visuospatial topographic representation in other cortical areas

Topographic representation of visual fields has been suggested in extrastriate cortical areas that are connected to the dorsolateral PFC and are involved in visuospatial information processing. For example, in areas 7a and PO, which are connected to the dorsolateral PFC, visual fields appear to be topographically organized because different regions of area 7a are topographically connected to different regions of area PO, which are in turn connected topographically to different regions of V1 and V2 that are associated with different portions of the visual field (Colby et al. 1988). Neuronal activities of area 7a during ODR performance are very similar to those of the dorsolateral PFC and are involved in visuospatial working memory processes (Chafee and Goldman-Rakic 1998). Therefore it is likely that the dorsolateral PFC is topographically connected to the posterior parietal area, as noted by Goldman-Rakic (1988). The posterior parietal area might also contain a memory map that is interconnected and interacts with the memory map of the dorsolateral PFC for representing visuospatial working memory processes; this problem should be examined in further studies.

In their study of the activity of single neurons, Suzuki and Azuma (1983) demonstrated that visual fields for visual perception/attention are topographically represented on caudal lateral regions of the PFC (this region seems to involve part of rostral area 8 and part of caudal area 46). Based on their findings, more peripheral visual fields are represented in a more medial region of the caudal lateral area of the PFC, and more central visual fields are represented in more lateral regions. In the case of a memory map, more peripheral memory fields are represented in more medial regions of the dorsal convexity of the PFC, and more central memory fields are represented in more lateral regions. Therefore the memory field represented in the dorsolateral PFC appears to be a mirror image of the representation of the visual field in the caudal lateral region of the PFC. Furthermore, in the FEF, which occupies part of area 8, the amplitude of the saccade evoked by electrical stimulation is represented topographically, with large and small saccades represented medially and laterally, respectively (Bruce et al. 1985). This representation also appears to be a mirror image of the representation of eccentricity in the memory map in the dorsolateral PFC. Since the dorsolateral PFC is connected to area 8 (Arikuni et al. 1980, 1988; Watanabe-Sawaguchi et al. 1991), it is likely that these two regions are connected topographically, and information is also processed topographically in these regions. Mirror-image representation of the visual field occurs in some visual cortical areas (e.g., V1, V2, and V3) that are adjacent to each other, and mirror-image somatic representation is present in somatosensory cortical areas, such as 3a and 3b (Kaas 1983, 1987, 1989, 1993). A similar principle of mirror-image representation may be present in areas of the dorsolateral PFC.

Topographic modular representation of working memory in the PFC

The connections between the dorsolateral PFC and other cortical areas, including the posterior parietal areas and area 8, are organized modularly; terminal fields as well as cells at the origin form a vertically arranged modular organization called a “module” or “column” (Arikuni et al. 1988; Bugbee and Goldman-Rakic 1983; Goldman-Rakic 1984; Goldman-Rakic and Schwartz 1982; Watanabe-Sawaguchi et al. 1991). The width of the corticocortical columns of the PFC has been estimated to be 400–1200 μm, with a median of ~700 μm in rhesus monkeys (Bugbee and Goldman-Rakic 1983). Columnar/modular organization has also been found for intrinsic connections within the dorsolateral PFC (Levitt et al. 1993; Pucak et al. 1996), and functional columns have been visualized in our preliminary studies with optical recording techniques with a voltage-sensitive dye in brain slices of the monkey dorsolateral PFC (Nakamura and Sawaguchi 1995; Sawaguchi 1996). Considering these and the present findings, it is likely that visuospatial memoranda are represented in different modules/columns that are arranged topographically in the dorsolateral PFC. The different modules may be connected to different modules of other cortical areas, such as posterior parietal areas and area 8, that are involved in visuospatial functions. Different channels formed by these different modules may be involved in a series of visuospatial working memory processes for different parts of the visual field.

Thus memoranda for visuospatial information appear to be represented as a memory map consisting of modules/columns arranged topographically in the dorsolateral PFC. However, the region examined as well as the target locations presented were limited, and we did not explore the depth of the sulcus. In addition, our method using MUS injection is limited with regard to fine-grained mapping. Therefore further studies with various techniques, such as optical imaging in vivo, are required to reveal the entire detailed memory map and to examine whether or not there are multiple memory maps within the PFC. Nevertheless, the present findings enable us to conclude that a specific site in the dorsolateral PFC is involved in the mnemonic function/process for a specific visuospatial coordinate, which should be important for understanding the functional organization of the PFC for representing visuospatial working memory processes.

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