Behavioral/Cognitive

Human Sensorimotor Cortex Represents Conflicting Visuomotor Mappings

Kenji Ogawa1 and Hiroshi Imamizu1,2

1Cognitive Mechanisms Laboratories, Advanced Telecommunications Research Institute International, Keihanna Science City, Kyoto 619-0288, Japan, and 2Center for Information and Neural Networks, National Institute of Information and Communications Technology and Osaka University, Suita, Osaka 565-0871, Japan

Behavioral studies have shown that humans can adapt to conflicting sensorimotor mappings that cause interference after intensive training. While previous research works indicate the involvement of distinct brain regions for different types of motor learning (e.g., kinematics vs dynamics), the neural mechanisms underlying joint adaptation to conflicting mappings within the same type of perturbation (e.g., different angles of visuomotor rotation) remain unclear. To reveal the neural substrates that represent multiple sensorimotor mappings, we examined whether different mappings could be classified with multivoxel activity patterns of functional magnetic resonance imaging data. Participants simultaneously adapted to opposite rotational perturbations (+90° and −90°) during visuomotor tracking. To dissociate differences in movement kinematics with rotation types, we used two distinct patterns of target motion and tested generalization of the classifier between different combinations of rotation and motion types. Results showed that the rotation types were classified significantly above chance using activities in the primary sensorimotor cortex and the supplementary motor area, despite no significant difference in averaged signal amplitudes within the region. In contrast, low-level sensorimotor components, including tracking error and movement speed, were best classified using activities of the early visual cortex. Our results reveal that the sensorimotor cortex represents different visuomotor mappings, which permits joint learning and switching between conflicting sensorimotor skills.

Introduction

Humans can concurrently learn and switch among various motor skills. There are two kinds of multiple-skill acquisition: one is learning the different types of perturbations, such as kinematics versus dynamics transformation, which involves independent processes in the sense that joint learning of two separate skills does not cause interference (Krakauer et al., 1999). The other type is adaptation to different parameters within the same type of perturbation, such as varying angles of visuomotor rotation, which sometimes induces conflicting interference (Wigmore et al., 2002). For example, people do not confuse riding a bike with driving a car, but they have difficulty in switching between driving on the left and on the right side of the road in different countries. Related to the latter type of adaptation, behavioral experiments have shown successful learning for conflicting force fields (Wada et al., 2003; Osu et al., 2004; Nozaki et al., 2006; Howard et al., 2008) or opposite visuomotor rotations (Cunningham and Welch, 1994; Gahramani and Wolpert, 1997; Imamizu et al., 2007b; Choi et al., 2008) after intensive training. Another study showed that exposure to various angles of visuomotor rotation induces structural learning (Braun et al., 2009). Previous literature revealed the involvement of separate brain regions in the learning of different types of perturbation (Imamizu et al., 2003; Krakauer et al., 2004; Girgenrath et al., 2008 for human neuroimaging; Rabe et al., 2009 for a lesion study), which is consistent with behavioral evidence of independent learning as well as the computational theory (Wolpert and Kawato, 1998). In contrast, the neural mechanisms underlying joint adaptation to the multiple parameters within the same types of perturbation remain unclear. To reveal the neural substrates that represent multiple sensorimotor mappings, we used functional magnetic resonance imaging (fMRI) to measure brain activities related to conflicting visuomotor mappings. However, the conventional univoxel fMRI analysis evaluates overall increases in activation of specific regions and would not be sensitive to distributed representations contained in patterns of voxel activity. To overcome this issue, this study used multivoxel pattern analysis (Haynes and Rees, 2005; Kamitani and Tong, 2005; Norman et al., 2006) for examining whether different mappings could be classified with distributed patterns of neural activities.

Using visuomotor tracking, participants simultaneously adapted to two types of visuomotor rotation (i.e., dual adaptation task) in which the cursor position of a joystick was rotated by +90° or −90° around the starting position. After adaptation, we classified the two rotation types during the tracking with the multivariate activity pattern of fMRI. One possible concern is that tracking under different rotation types could be accompa-
Materials and Methods

Subjects. Subjects were seven male volunteers with a mean age of 30.0 years (range, 22–49 years). All subjects were right-handed, as assessed by a modified version of the Edinburgh Handedness Inventory (Oldfield, 1971). Written informed consent was obtained from all subjects in accordance with the Declaration of Helsinki. The experimental protocol received approval from the local ethics committee.

Task procedures. We used a continuous visuomotor tracking task (Imamizu et al., 2000, 2003, 2004, 2007a): subjects moved a small crosshair cursor on the screen with a joystick and continuously tracked a moving target. After a starting message shown for 2 s, target and cursor were presented on the screen. The target was a small green circle on a dark background, which moved around within a square subtending horizontal and vertical visual angles of −14°. The cursor was a small yellow cross. We introduced a rotational perturbation between cursor positions and a joystick angle of 90° around the screen center, in either clockwise or counterclockwise direction. During the task block of 21 s, subjects continuously tracked the moving target. A fixation dot was displayed for 4 s during intertrial intervals (ITI) (Fig. 1A). After a practice session to become familiarized with the scanner environment, the subject underwent five sessions, with each consisting of 24 blocks. One session lasted −11 min. The block order was counterbalanced as A-B-B-A-A-B-B-A… with A and B pseudorandomly assigned the two rotation types across sessions. Unlike previous studies (Osu et al., 2004), we did not provide visual or auditory cues for different rotation types to equate sensory inputs between conditions.

Stimuli were presented on a liquid crystal display and projected onto a custom-made viewing screen. Subjects lay supine in the scanner and viewed the screen via a mirror, being unable to see their hand throughout this task. They used their right index and middle fingers to control the joystick, with the right upper arm immobilized using foam pads to minimize body motions. A custom-made bite bar was used for all participants to prevent head motions for all the scanning sessions. The amount of subjects’ head motion, which was estimated from alignment of functional images (see below), ranged from 0.07–2.0 mm, with a median of 0.36 mm within a session.

Functional localizer scans. On a separate day from the main rotational experiment, an independent localizer scan was performed for all participants to define sensorimotor areas relevant to visuomotor tracking. The task was the same as in the main experiment except that it had a longer baseline period to reliably detect increased fMRI responses related to visuomotor tracking, and that subjects used a normal joystick without rotation. One session lasted ~12 min and had 24 task blocks of 15 s, interleaved with 15 s rest periods using a fixation display.

Target trajectory. Stimuli were generated using Psychophysics Toolbox (http://www.psychtoolbox.org; Brainard, 1997) for MATLAB (MathWorks). The x- and y-components of the target trajectory were the sums of sinusoids whose amplitudes and frequencies were pseudorandomly determined using the same procedure as in our previous studies (Imamizu et al., 2000, 2003, 2004, 2007a). Since these parameters were determined pseudorandomly for each run, the target paths differ across sessions and among subjects. In addition, two different patterns of target trajectory were created to dissociate low-level movement kinematics or reafferent sensory inputs with the rotational perturbations. Successful classification of rotation types would support the existence of neural substrates representing conflicting visuomotor mappings.

Figure 1. A, Time course of experiment. After display of start message for 2 s, target (circle) and cursor (cross) were presented on the screen. Subjects continuously tracked a moving target with cursor using joystick for 21 s, with rotational perturbations of ±90° or −90°. Task blocks were interleaved with rest periods of 4 s. B, Examples of target trajectory under different rotation types and velocity types in single block. C, Examples of horizontal positions of target with those of cursor within single block of first (left) and last (right) practice sessions.
Behavioral training. Before the fMRI scanning, subjects underwent 9–16 sessions of training outside the MRI scanner for 3 or 4 d. The training session is the same as those used for fMRI scanning, including the same rotation perturbations, except that ITI is shorter (2 s). One session lasted ~10 min. One subject (author K.O.) did not participate in the training sessions because he had already undergone sufficient practice during the preliminary experiment.

Behavioral analysis. Cartesian coordinates of the on-screen target and cursor positions were recorded at 60 Hz. We then calculated the following measures separately for x- and y-components: (1) tracking error defined as a Euclidean distance between target position and cursor position, (2) distance of cursor trajectory, (3) correlation coefficient between target and cursor trajectories, and (4) correlation coefficient between x- and y-components. These behavioral measures were calculated for each block, normalized as a z-score among all trials within subjects and then compared both between the two conditions of rotation (+90° vs. −90°) and between those of velocity (fast vs. slow). The power spectrum of cursor velocity was calculated with fast Fourier transform to measure the intermittency, which evaluates the frequency of on-line corrections based on visual feedback in manual tracking (Miall, 1996; Foulkes and Miall, 2000).

MRi acquisition. A 3 T Siemens Trio scanner with a 12-channel head coil was used to perform T2*-weighted echo planar imaging (EPI). A total of 222 and 246 scans were acquired for the main and localizer sessions, respectively, with a gradient echo EPI sequence. The first four scans were discarded to allow for T1 equilibration. Scanning parameters were TR, 3000 ms; TE, 30 ms; FA, 80°; FOV, 192 × 192 mm; matrix, 64 × 64; 50 axial slices; and slice thickness, 3 mm without gap. T1-weighted anatomical imaging with an MP-RAGE sequence was performed with the following parameters: TR, 2250 ms; TE, 3.06 ms; FA, 9°; FOV, 256 × 256 mm; matrix, 256 × 256; 192 axial slices; and slice thickness, 1 mm without gap.

Processing of fMRI data. Image preprocessing was performed using SPM8 software (Wellcome Department of Cognitive Neurology; http://www.fil.ion.ucl.ac.uk/spm). All functional images were first realigned to adjust for motion-related artifacts. The realigned images were then spatially normalized with the Montreal Neurological Institute (MNI) template, based on affine and nonlinear registration of coregistered T1-weighted anatomical images (normalization procedure of SPM), and resampled into 3 mm cube voxels with sinc interpolation. All images were spatially smoothed using a Gaussian kernel of 8 × 8 × 8 mm full-width at half-maximum. The smoothing was not performed except for the localizer scans, as this could blur fine-grained information contained in multivoxel activity (Mur et al., 2009).

Using the general linear model, the 24 blocks per session were modeled as separate 24 box-car regressors that were convolved with a canonical hemodynamic response function. Our previous study using the same tracking task showed that switching of different visuomotor perturbations takes ~5 s after block onset (Imamizu et al., 2004). We thus removed the initial 6 s, corresponding to two fMRI scanings, to remove transient blood oxygenation level-dependent activity at task onsets as associated with task switching or increased tracking error, and thus box-car length was 15 s covering the latter part of a 21 s block. The behavioral analysis was also conducted by removing the first 6 s from the realignment stage were modeled as confounding covariates. Low-frequency noise was removed using a high-pass filter with a cutoff period of 128 s, and serial correlations among scans were estimated with an autoregressive model implemented in SPM8. This analysis yielded 24 independently estimated parameters (β values) per session for each individual voxel, which were subsequently used as inputs for decoding analysis.

Definition of regions of interest. The regions of interest (ROIs) were defined for each subject, both functionally with activations of localizer scans (Fig. 2) and anatomically with the automated anatomical labeling (AAL) toolbox (Tzourio-Mazoyer et al., 2002) or Brodmann areas included in the MRICro software (http://www.mricro.com). First, the localizer scan was used to identify the activated areas that showed significantly higher response during a task epoch compared with the baseline period with an uncorrected threshold of p < 0.01 at voxel level. The primary motor and lateral premotor areas (Mot) and the somatosensory area (Som) were then anatomically masked with the precentral and postcentral cortices of AAL, respectively. The supplementary motor area (SMA) was also defined using AAL mask. The early visual area was anatomically masked within Brodmann areas 17 and 18. In the posterior parietal cortex, we selected the medial intraparietal cortex (miPFC), which is related to visually guided hand movements (Ogawa and Inui, 2012). The parietal ROI was masked both with miPFC having a 15 mm radius sphere around the peak average of previously reported coordinates of [−29, −51, 58] (Greffkes et al., 2004; Prado et al., 2005) and anatomically with the superior parietal lobule (SPL). The anterior and posterior parts of the cerebellum were anatomically defined based on the primary fissure. As a control ROI, we chose voxels outside of the brain centered at [57, 60, 57] with a radius of 15 mm (nonbrain). Several areas of one subject did not show significant activations in the localizer scan, and thus ROIs were based on activations of group-level results of other subjects (n = 6) only for this participant. Table 1 describes mean (SD) MNI coordinates and the number of voxels of ROIs.

Mass univariate analysis. For the conventional univariate analysis of individual voxels, we directly compared two conditions for both rotations (+90° vs. −90°) and velocity (fast vs. slow). Contrast images of each...
subject, generated using a fixed-effects model, were taken into the group analysis using a random-effects model of a two-way ANOVA with rotation and velocity types as within-subject factors. Activation was reported with a lenient threshold of $p < 0.01$ uncorrected for multiple comparisons at voxel level with an extent threshold of 10 voxels. The ROI analysis was additionally performed by comparing average parameter estimates ($\beta$ values) between the two conditions of rotation types as well as velocity types.

**Multivoxel pattern analysis.** The multivariate classification analysis of fMRI was performed with a binary classifier based on linear support vector machine (SVM) implemented in LIBSVM (http://www.csie.ntu.edu.tw/~cjlin/libsvm/), with default parameters (a fixed regularization parameter $C = 1$). Parameter estimates ($\beta$ values) of each trial of voxels within ROIs were used as inputs to the classifier.

We conducted two types of classification analysis. First, we classified the rotation types with cross-validation between different velocity types. The decoder was first trained to classify fast and slow target with target trials, and the same decoder was then used to classify fast and slow with 90° rotation and slow target. The same decoder was then used to classify fast and slow with 90° rotation and fast target with 90° trials. Second, we used different combinations of rotation types and velocity types for training and testing datasets: the classifier was first trained to discriminate the trials with rotation of 90° and fast target from those of 90° rotation and slow target. The same decoder was then tested to see if it could classify the trials of 90° rotation and fast target separately from those of 90° rotation and slow target. We also conducted classification in the reverse direction. In this analysis, manipulation of rotation types and velocity types was orthogonal to each other, and thus classification accuracy of rotation was the opposite to that of velocity (i.e., the accuracy of rotation equals to 100% minus the accuracy of velocity; Fig. 3B). This was done to explicitly dissociate low-level movement kinematics or reafferent sensory inputs associated with rotation types, since all types of behavioral differences including tracking error and movement distance were larger in velocity types compared with rotation types (see Results, Behavioral performance). Such a “cross-classification,” a cross-validation between trials with different sets of tasks or stimuli, has been previously used to investigate the similarity or invariance of neural representations by testing the generalization of a classifier between different conditions or modalities (Etzel et al., 2008; Knops et al., 2009; Stokes et al., 2009; Meyer et al., 2010; Oosterveld et al., 2010; Gallivan et al., 2011a; Kahnt et al., 2011; Ogawa and Imamizu, 2011, 2012; Corradi-Dell’Acqua et al., 2011). A cross-validation was conducted 10 times (5 sessions $\times$ 2 combinations) and then the average accuracy was estimated.

Tests of statistical significance for classification accuracy were conducted in both parametric and nonparametric methods. First, a two-tailed test was used to determine whether the observed decoding accuracy was significantly higher than chance ($50\%$) with intersubject difference treated as a random factor ($df = 6$). To control for the problem of multiple comparisons, we applied the Holm–Bonferroni procedure (Holm, 1979) based on the number of ROIs. Second, we conducted a randomization test that was identical to the decoding analysis described previously, except that we randomly shuffled correspondence between the fMRI activations and the condition labels before training the classifier within each subject and obtained average decoding accuracy across participants. This procedure was separately repeated 1000 times for each ROI by reshuffling the labels each time, which created an empirical distribution of decoding accuracy under a null hypothesis of chance. A significant above chance accuracy of $p < 0.05$ was assumed if the decoding accuracy with correct labeling exceeded the 95th percentile of the null distribution.

Complementary to the a priori ROI analysis, we additionally conducted a volume-based “searchlight” analysis (Kriegeskorte et al., 2006). Cross-classification with different pairs of rotation and velocity types (Fig. 3B) was performed using multivoxel activation patterns within a 9 mm radius sphere (searchlight) containing at least 100 voxels (maximum of 123 voxels). The searchlight moved over the gray matter of the whole brain, and the average classification accuracy for each searchlight with leave-one-session-out cross-validation was assigned to the sphere’s center voxel. The resulting map of the decoding accuracy was averaged over subjects. Results were first thresholded at voxel level using a randomization test as in the ROI analysis. We conducted the same searchlight analysis by randomly shuffling the correspondence between the voxel activities and the condition labels, and estimated the average accuracy across participants. This procedure, which was repeated reshuffling the labels each time, created an empirical distribution of decoding accuracy under a null hypothesis of chance, which was then collapsed over all searchlights (the number of searchlights was 35,512, and so the number of data in the null distribution was 3,551,200). For each searchlight (i.e., each center voxel), a significant above-chance accuracy of $p < 0.0001$ was assumed if the observed accuracy with correct labeling exceeded the 99.99th percentile of the null distribution. To correct for multiple comparisons, a cluster-level threshold was further applied with a randomization test (Nichols and Holmes, 2002; Oosterveld et al., 2010) using the same permutation distribution generated above. We obtained a maximum cluster size (i.e., the number of voxels) for each randomization sample at the same voxel-level threshold, which yields an empirical null distribution of a maximum cluster extent. Clusters were reported as significant at a cluster level of $p < 0.01$ if the observed number of voxels within the cluster with correct labeling exceeded the 99th percentile of this null distribution.

**Additional experiment with normal joystick.** To further confirm that the classification result of rotation types was not confounded with differences in movement kinematics, we conducted another experiment with five of the seven subjects who participated in the main experiment. This time subjects used a normal (not rotated) joystick to track a target whose motions followed their own joystick trajectory of the previous rotational experiments. The task design and number of trials as well as sessions per subject were the same as those in the rotational experiment. We hypothesized that if differences in movement patterns between rotational conditions had contributed to successful classification of rotations, significant classification accuracy could also be obtained in this control experiment.

**Eye-movement recording.** Our subjects freely made eye movements during the tracking, which could be a potential confound. To exclude the possibility that differences in eye movements contributed to the successful classification of rotation types, we measured the eye positions during the rotational tracking outside the fMRI scanner after the main fMRI experiment. The horizontal and vertical eye positions of four among seven subjects who participated in the main experiment were measured at 60 Hz with the Tobii X120 infrared recording system (Tobii Technology; http://www.tobii.com). The task sequences were the same as those of the fMRI experiment except for a shorter trial duration (12 s) to minimize eye blinks; five sessions were conducted per participant. We then ran the classification analysis based on the time series of the eye positions as well as the power spectrum of the eye movement velocity.
with the same procedure as multivoxel pattern analysis (MVPA) using different combinations of rotation types and velocity types (Fig. 3B).

**Results**

**Behavioral performance**

Over the training sessions, subjects gradually learned two rotational types, as indicated by the decrease in tracking errors as the number of sessions proceeded (Fig. 4A). We compared the average tracking errors of the first and the last three sessions, respectively, for each subject, and found a significant decrease (paired t test, t(5) = 6.49, p = 0.001; Fig. 4B). We next analyzed the maximal cross-correlation between the trajectories of the target and the cursor averaged during the first and last three sessions. This revealed a significant increase in correlation coefficients, indicating that the cursor followed the target trajectory more accurately in the later sessions (paired t test, t(5) = 4.55, p < 0.01; Fig. 4C).

The time series of tracking errors within a 21-s block was compared between rotation and velocity types (Fig. 5A). Because a transient increase in errors was found after block onset, behavioral and fMRI data were used only after 6 s of trial onset (Fig. 5A, gray areas) for later analysis. We also compared the power spectrum of the cursor velocity between the two conditions (Fig. 5B), which showed the differences in the cursor movement frequency or the intermittency of the feedback control. The figures show that differences in both tracking errors and power spectrum were larger between velocity types compared with rotations. For quantitative comparison, we calculated various behavioral measures for cursor trajectory and compared the normalized absolute differences between the two conditions of rotation and velocity types (see Materials and Methods, Behavioral analysis for details; Fig. 5C). We then compared the differences between the rotation and velocity types and found significantly larger differences between the two velocity types compared with the rotation types for measures including tracking error and movement distance, while no measure showed significantly greater differences in the reverse direction. These behavioral results indicate that the kinematic differences are larger in the two velocity types compared with those of the rotation types, which ensures that a higher decoding accuracy of rotation types than velocity types later found in MVPA cannot be due to kinematic differences.

**Mass univariate analysis**

We directly compared two conditions of both rotation (+90° vs −90°) and velocity (fast vs slow), using the traditional univariate analysis of single voxels. A significant main effect of velocity types was observed in the early visual areas bilaterally at a lenient threshold of p < 0.01 uncorrected for multiple comparisons (Fig. 6A). No significant main effect of rotation types or interactions was found at the same threshold. In addition, the ROI analysis comparing averaged parameter estimates (β values) between the two conditions of rotation and velocity within each ROI revealed no significant difference in either rotation or velocity comparison (Fig. 6B).

**MVPA**

First, we classified the rotation types with the cross-validation between different velocity types to test whether the decoder trained to classify two rotation types in one motion pattern could be generalized to the other (Fig. 3A). While the statistical results were described based on the corrected threshold unless otherwise mentioned, we included both the corrected and uncorrected significance in Figure 7 for completeness. The results showed significant above-chance classification accuracy of rotation types in Mot (62.2%, t(6) = 10.97, p < 0.001), Som (67.3%, t(6) = 4.14, p < 0.01), SMA (62.3%, t(6) = 6.10, p < 0.001), early visual area, Vis (60.6%, t(6) = 4.80, p < 0.05), anterior part of cerebellum, CBLa (61.5%, t(6) = 3.34, p < 0.05), and posterior part of cerebellum, CBlp (61.3%, t(6) = 4.46, p < 0.005). The successful classification of velocity types between different rotation types was found in Vis (78.9%, t(6) = 11.33, p < 0.001) (Fig. 7A).

Since we found small differences in the behavioral measures between two rotation types and the velocity types (Fig. 5C), the successful classification of rotation types might reflect differences in the low-level kinematics or the resultant visual feedback of
cursor motions. The higher than chance decoding accuracy for rotations together with the velocity types in Vis suggests that slight differences in kinematics could also contribute to the classification of rotation and velocity types. To exclude that possibility, we then performed “cross-classification” with different combinations of rotation and velocity types to explicitly dissociate the movement kinematics with rotation types (Fig. 3B). If the kinematics contributed to successful classification of rotation types, the accuracies for velocity types should exceed those of rotation types in this analysis, because the differences in all the behavioral measures were larger among the velocity types compared with rotation types (Fig. 5). Significant decoding accuracy of rotation was found for Mot (59.2%, $t_{(s)} = 9.21$, $p < 0.001$), Som (62.1%, $t_{(s)} = 3.12$, $p < 0.05$), SMA (57.6%, $t_{(s)} = 2.94$, $p < 0.05$), and CBLa (57.5%, $t_{(s)} = 2.78$, $p < 0.05$), while that of velocity was found for Vis (71.9%, $t_{(s)} = 6.94$, $p < 0.001$) (Fig. 7B). Among these ROIs, Mot and Vis showed significant above-chance accuracy after correction for multiple comparisons, and the results of each individual subject (denoted by small dots in Fig. 7B) show high consistency. We additionally conducted a nonparametric randomization test and found that it generally produced a more lenient threshold for $p$ values compared with the parametric $t$ test (similar results were reported by Gallivan et al., 2011b). This confirmed that all of the significant above-chance accuracies found by the $t$ test were also significant by the randomization test.

The same classification was also conducted by equating the number of voxels across ROIs to eliminate differences in feature size. The voxels in each ROI were selected in the order of highest $t$ value of the localizer scan, which was based on the univariate analysis, until the number of voxels reached 400 for each subject (Pereira et al., 2009). This replicated the previous results: significant decoding accuracy of rotation for Mot (56.2%, $t_{(s)} = 4.79$, $p < 0.05$), Som (60.3%, $t_{(s)} = 2.74$, $p < 0.05$), and SMA (57.3%, $t_{(s)} = 2.45$, $p < 0.05$), and that of velocity for Vis (70.0%, $t_{(s)} = 5.59$, $p < 0.005$) with uncorrected $p$ values.

Next, the same classification was performed with averaged $\beta$ values of all voxels within an ROI as inputs. No significantly higher than chance accuracy was found in any ROI, which indicates that decoding was based on differences in distributed spatial patterns of multiple voxels, rather than overall amplitude differences within the ROI (Fig. 7C). This is also supported by the voxel bias map, which displays the averaged weights (positive or negative) of the classifier across 10 folds (i.e., iterations) in the cross-validation for individual voxels within ROIs; this map showed intermingled patterns of voxels with two rotational types (+90 vs −90 biased voxels) within ROIs in all subjects (Fig. 8A for a single typical subject). In addition, the bias patterns between different pairs of participants showed low correlations around zero (Fig. 8B), which indicates idiosyncratic patterns of weights specific to each subject.

Finally, the searchlight map revealed the voxels for higher accuracy of rotation types in the sensorimotor cortex as well as in the SMA (Fig. 9, red). These areas closely correspond to the activated regions in the localizer scans that were conducted separately (Fig. 2A). We also found the voxels for higher accuracy of velocity types mostly located in the early visual areas (Fig. 9, blue). These are largely consistent with the results of the previous ROI analysis.

**Classification with behavioral measures**

To further validate that successful classification of rotation types is not due to differences in low-level movement kinematics, we conducted multivariate classification analysis using the various behavioral measures. In this analysis, all of the observed behavioral measures (see Materials and Methods, Behavioral performance) were used separately or jointly as features (Fig. 10A), and classification of rotation types versus velocity types was conducted in the same way as the fMRI–MVPA (Fig. 3B). The time series of tracking error as well as the power spectrum of cursor trajectory was also used as a feature vector for classification (Fig. 10B). These analyses revealed significantly greater classification accuracy only between velocity types compared with rotations. We further conducted the same classification analysis using the

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**Figure 5.** Behavioral results during fMRI scanning. **A**, Time series of mean tracking error within block. Two rotation and velocity types were compared in left and right figures, respectively. Trial onset of 0.5 (gray areas) was removed for later analysis. Shaded region with colors denoted 5D across subjects. **B**, Power spectrum of cursor trajectory averaged over trials and two dimensions during fMRI scanning. Two rotation and velocity types were compared in left and right figure, respectively. **C**, The comparisons of behavioral measures between rotation and velocity types.
time series as well as the power spectrum of the eye movements, which again showed a higher accuracy for the velocity types compared with the rotation types (Fig. 10C). These results indicate that higher classification accuracy for rotation over velocity in MVPA (Fig. 7B) cannot be attributed to the differences in the movement or the sensory reafferents associated with rotational perturbations as well as in eye movements.

Additional experiment with normal joystick

We additionally conducted the same MVPA using different combinations of rotation types and velocity types (Fig. 3B) when subjects used a normal (not rotated) joystick to track a target in a control experiment. The averaged (SD) maximum cross-correlations between the target and cursor trajectory of the x- and y-directions were 0.78 (0.05) and 0.81 (0.04), respectively. This indicates that all of the subjects accurately followed the target trajectory. A significant decrease in classification accuracies for rotation types compared with those of the main rotational experiment was observed in Mot (paired t test; \( t_{(4)} = 5.38, p < 0.01 \)), Som (\( t_{(4)} = 4.11, p < 0.05 \)), and CBLa (\( t_{(4)} = 3.30, p < 0.05 \)), and a marginal significance was found in SMA (\( t_{(4)} = 2.41, p = 0.07 \)), with nonsignificant differences with the chance level in the control experiment (all \( p > 0.5 \)) (Fig. 11). In a post experimental debriefing, none of the subjects could discriminate whether a target motion was from their cursor trajectory of the +90° or −90° condition in the previous rotational experiment, while they could notice fast versus slow target trajectories. This indicates that movement kinematics were more similar between the two rotation types than between the velocity types, which is consistent with the results of behavioral analysis (Figs. 5, 10).

Discussion

This study aimed to reveal neural substrates that represent conflicting visuomotor mappings. After intensive training, the participants jointly adapted to the two rotational perturbations, which is indicated by the behavioral measures (Fig. 4). The fMRI decoding showed that rotation types were successfully classified with multivariate activity patterns in the sensorimotor cortex when trials with different types of rotation and velocity were combined and cross-validated in the ROI analysis (Fig. 7B). Searchlight analysis also revealed a significant above-chance accuracy of rotation types in the sensorimotor cortex and the SMA (Fig. 9). Our findings reveal that the sensorimotor cortex represents different mappings, which subserves simultaneous adaptation of multiple visuomotor correspondences.

This successful classification cannot be due to differences in low-level movement kinematics or sensory reafferents associated
with the rotational perturbations for the following two reasons. First, the differences in the various behavioral measures, including tracking error, movement distances, and correlation of target and cursor motions, as well as power spectrum, are larger between the velocity type compared with the rotation type (Fig. 5). The classification using these behavioral measures as well as eye movements, conducted in the same way as the fMRI-MVPA, further showed significantly higher accuracies of velocity types over rotation types (Fig. 10). Second, the additional experiment, in which subjects tracked their joystick trajectory with a normal joystick, showed a significant decrease in accuracy for rotation types in the same ROIs with a chance-level performance (Fig. 11). These results indicate that successful decoding of rotation types is not confounded with movement kinematics. In contrast, low-level sensorimotor components, which were associated with different velocity types including tracking error and movement speed, were best classified using activities of the early visual cortex (Figs. 7, 9). We also found a significantly higher than chance decoding accuracy of velocity types as well as rotation types in the sensorimotor cortex and the cerebellum when classification of velocity types was cross-validated between different rotation types (Fig. 7A). This indicates that low-level sensorimotor components were also represented in these areas, together with the rotational types. Previous studies with MVPA or fMRI adaptation (Grill-Spector and Malach, 2001) revealed the directional selectivity of hand movements in the primary motor cortex (Eisenberg et al., 2010, 2011; Fabbri et al., 2010; Ogawa and Inui, 2012). Eisenberg et al. (2011) introduced a single rotational perturbation (45°) between hand and cursor movements to dissociate movement directions in the visual and motor coordinates. In contrast, our current study introduced the two conflicting rotational perturbations between hand and cursor movements (+90 or −90°), requiring participants to move their hands in an opposite direction depending on the rotation types. We then decoded the rotational directions of the visuomotor perturbation that are independent of the movement directions. Our study indicates, for the first time, that multiple visuomotor mappings can be decoded with distributed neural activity patterns of fMRI. Previous human neuroimaging studies revealed that separate brain regions are involved in the learning of different types of perturbation (Imamizu et al., 2003; Krakauer et al., 2004; Girgenrath et al., 2008), which indicates that the size of these separate modules are larger than fMRI individual-voxel resolution. In contrast, our results showed that parametric differences in the same type of perturbation cannot be discriminated with the mass univoxel analysis (Fig. 6) but could be revealed using MVPA. The voxel bias map showed intermingled patterns of voxels with two rotational types idiosyncratic to each participant (Fig. 8). The previous studies showed that orientation columns in the early visual cortex could be discriminated using MVPA that is beyond fMRI voxel resolution (Haynes and Rees, 2005; Kamitani and Tong, 2005). Our results suggest that parametric differences within the same perturbation are represented more closely and intermixed at a resolution below that of fMRI voxels, compared with different perturbations in separate areas that are larger than voxel size. This discrepancy may correspond to the existence of behavioral interference: joint learning of the different types of perturbations, such as kinematics versus dynamics, does not cause interference (Krakauer et al., 1999), while adaptation to different parameters within the same type of perturbation, like that in the current...
study, induces conflicting interference (Wigmore et al., 2002). This interference could be caused by the conflicting use of the same neural representations, which are more closely located for multiple parameteric mapping within the same type rather than different types of perturbations. Noted that the possible information source of MVPA remains unclear with some recent controversies (Freeman et al., 2011; Beckett et al., 2012). The decoding accuracy of MVPA is considered to be dependent on the spatial patterns of distinct neuronal populations and/or accompanying vasculature units together with the spatial resolution of fMRI (Bartels et al., 2008; Gardner, 2010). Our results are also in line with recent papers showing the effectiveness of MVPA in motor systems by successfully decoding multiple actions (Dinstein et al., 2008; Gallivan et al., 2011a,b), representations of individual fingers (Wiestler et al., 2011; Diedrichsen et al., 2013), or movement directions (Eisenberg et al., 2010, Ogawa and Inui, 2011, 2012), which are not possible with the conventional univoxel analysis.

There are two possibilities for the neural mechanisms underlying the joint adaptation of the multiple parameters within the same types of perturbation. One is that a single component that represents the visuomotor correspondence continuously adapts and readapts to different mappings depending on the current environment, which regards switching as recalibration between the visual and motor frames of reference. The other is that multiple mappings have distinct substrates, each of which represents a specific visuomotor correspondence (Lee and Schweighöfer, 2009). In our current study, it remains unclear whether the motor system serially recalibrates a single general model or acquires multiple models specific to each mapping, since both possibilities induce specific distributed activation patterns of multiple voxels for decoding. Further studies are needed to clarify whether the distinct mappings are represented with a single or multiple models.

Previous research identified the sensorimotor cortex and cerebellum as neural substrates of visuomotor learning (for review, see Shmuelof and Krakauer, 2011). Neurophysiological studies with nonhuman primates revealed changes in neuronal response reflecting learning of rotational perturbations in the primary motor cortex (M1) (Wise et al., 1998; Paz et al., 2003; Paz and Vaadia, 2004) as well as in the SMA (Padoa-Schioppa et al., 2004; Paz et al., 2005). A recent study using unit recording of nonhuman primates suggested that two opposing rotations cannot be represented with a single neuron in M1 (Zach et al., 2012). Their results support the existence of separate representations for different mapping, consistent with our current findings. Together with these previous unit recordings of nonhuman primates, our results indicate the existence of different neuronal populations in the motor-related region.

Human neuroimaging studies also indicated increased activations (Karni et al., 1995) as well as structural changes in M1 related to motor learning (Draganski et al., 2004; Landi et al., 2011). The recent studies using transcranial magnetic stimulation or transcranial direct current stimulation indicate the acquisition (Hunter et al., 2009) as well as storage (Hadipour-Niktarash et al., 2007; Galea et al., 2011) of sensorimotor skills in the human M1. The error-related increase in fMRI signal was also reported in the human sensorimotor areas when subjects learned visuomotor perturbations, which could be used as training signals for acquisition of internal models (Diedrichsen et al., 2005). These findings indicate the role of the sensorimotor cortex in representing motor skills, and they are generally consistent with the view that the primary motor cortex is not a low-level static controller of muscles but rather contains higher level representation of various movement parameters (for review, see Scott, 2003). Our results support this view, indicating that distinct visuomotor mappings are stored in the sensorimotor cortex.

Human neuroimaging has shown activities related to the acquisition of an internal model in the cerebellum (Imamizu et al., 2000, 2003; Seidler and Noll, 2008). Imamizu et al. (2003) showed that the learning of different types of sensorimotor perturbations (rotation and velocity) is subserved with the acquisition of multiple internal models in the cerebellum. Our study revealed significant above-chance accuracy in the anterior cerebellum in the ROI analysis, which supports the existence of multiple internal models in the cerebellum. However, since the accuracy in the cerebellum did not survive the threshold corrected for the multiple comparisons in the both the ROI and the searchlight analysis, we do not consider the cerebellum result to be robust and discuss it no further.

Previous studies indicated a role of the posterior parietal cortex (PPC) in visuomotor rotational learning (Inoue et al., 2000; Graydon et al., 2005). In contrast, we did not observe significant classification accuracy in the parietal cortex. While the previous research reported changes of activity in the midst of learning, our studies focused on activations when subjects sufficiently adapted to rotations after intensive practice. Our results indicate that the parietal cortex might be related to the early middle phases of a learning process, rather than long-term storing of motor memories. A recent computational model-based study also predicted that the site for learning is not in the PPC but in the connectivity between the PPC and the motor areas (Tanaka et al., 2009).

This study investigated opposite rotational perturbations that cause interference in joint learning. However, it is known that positive transfer of learning occurs when the rotational degrees of perturbations are close to each other (e.g., 30° or 45°) (Seidler and Noll, 2008). Future studies are needed to examine whether distinct visuomotor mappings exist in such cases, or if multivoxel pattern analysis can identify the mappings for close perturbations, if any. Extension of our studies to joint learning between various degrees of visuomotor rotations would give us important clues to understanding the modular units used for sensorimotor learning as well as the neural mechanisms used for positive and negative (interference) transfer of learning.

In summary, our findings indicate that the sensorimotor cortex represents conflicting visuomotor mappings, which subserves...
simultaneous adaptation of multiple sensorimotor mappings. We hypothesize that similar neural mechanisms underlie joint adaptation of opposing force fields. In future applications, decoding of a specific parameter in sensorimotor transformation could be used as a neurofeedback signal to achieve efficient structural learning with reduced interference (Braun et al., 2009), using a method recently applied to perceptual learning (Shibata et al., 2011).

References
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