INTRODUCTION

It is common knowledge that cortical representations in sensory-motor areas are not fixed but are continuously modified by experience (see Buonomano and Merzenich, 1998). When a lesion is made such that an adult animal loses sensation from a particular area, the region of the cortex innervated by the missing nerves loses inputs. However, after several weeks, this region becomes active again, being innervated by other axons adjacent to the lesion. Changes in cortical representation also occur as a result of training for tasks that produce specific, differential patterns of activity in cortical sectors. Skill learning or repetitive tactile stimulation expands digit representations in the somatosensory cortex of monkeys (Jenkins et al., 1990) and humans (Elbert et al., 1995), and in the human primary motor cortex (Karni et al., 1995, 1998). Reorganization is also known in the cerebellum regarding phylogenetically older sensory-motor parts. That is, a fractured tactile map reorganizes after deafferentation of a particular body part in adult rats (Shumway et al., 1999).

In the past decade, a number of studies have revealed that the cerebellum contributes to higher cognitive functions. Some of the first examples of a clearly cognitive task were reports of language processing and activation in the right lateral cerebellum (Petersen et al., 1989; Raichle et al., 1994). Since then, other studies (see Thach, 1996; Desmond and Fiez, 1998) have also pointed to cerebellar involvement in such cognitive tasks as working memory (Desmond et al., 1997) and problem solving (Grafman et al., 1992; Kim et al., 1994). Many studies (Desmond et al., 1997; Sakai et al., 1998; van Mier et al., 1998) have demonstrated that cognitive processing exists in the lateral part of the cerebellum. However, little is known about whether the organization of the lateral cerebellum and cerebro-cerebellar communication loop changes due to experience.

A recent positron emission tomography (PET) study on monkeys revealed that tool use activates a cerebro-cerebellar communication loop including the intraparietal sulcus (IPS) regions, the supplementary motor area (SMA), the premotor cortex, and the lateral cerebellum (Obayashi et al., 2001). Our previous study (Imamizu et al., 2000) demonstrated that an internal model for a novel tool (a rotated mouse whose cursor appeared in a position 120° rotated around the center of a screen) is acquired in lateral parts of the human cerebellum. Tamada et al. (1999) have shown that functional connectivity between the lateral parts and the premotor regions increased after acquisition of the internal model, suggesting that the internal model...
sends outputs to the premotor regions. We investigated cerebellar activity after subjects learned how to use two novel tools (the rotated mouse and a velocity mouse whose cursor velocity was proportional to the mouse position) (Imamizu et al., 1998, 2003). The brain locations activated while using the two different tools were spatially segregated with a small overlap, suggesting that multiple internal models are acquired in a modular fashion. Our study (Imamizu et al., 2004) suggested that the dorsolateral prefrontal cortex, the insula and parietal regions contribute to the switching and selection of multiple internal models as “central executives” or “responsibility estimators” in a MOdular Selection And Identification for Control (MOSAIC) model proposed for the learning and switching of internal models (Wolpert and Kawato, 1998a).

Here, using our experimental paradigm, we investigated the reorganization of internal models and changes in neural networks between the cerebellum and the frontal and parietal regions due to the acquisition of a new internal model. After long-term training with the two novel tools, subjects were briefly but intensively trained on the use of a third novel tool. Brain activities for the three types of tools were scanned before and after training for the third tool and compared.

A computer program mapping the joystick position to the cursor position makes the joysticks functionally different, and there is no difference in their form or fundamental manipulation. Although there might be some argument about whether we can consider them as entirely different “tools”, we can find common tools that have similar form but require a different skill. For example, scissors are similar to pliers in form and essential mechanism. However, pliers need a grip force in a direction parallel to the handle movements, while scissors need not only the grip force but also a force to antagonistically push the two edges. Each joystick in our experiment requires a different skill to control the cursor, depending on different mapping between the cursor movement and the joystick movement. Thus, our experimental paradigm can probe into how different skills for tool use are organized in the brain.

**METHODS**

**Subjects**

Fourteen neurologically normal subjects (five females and nine males from 21 to 39 years of age) participated in the experiments. All were right-handed according to the Edinburgh handedness inventory (Oldfield, 1971). Informed written consent was obtained from each subject. The test protocols were approved by the ATR ethics committee.

**Tasks**

Basic task designs are the same as those used in our previous studies (Imamizu et al., 2000, 2003, 2004). The subjects moved a small crosshair cursor on the screen with a joystick and continuously tracked a moving target. They controlled the joystick with their right hand. The target was a small white square on a dark background and moved around within a square area subtending horizontal and vertical visual angles of 7.33°. The x-and y-components of the target trajectory were sums of sinusoids whose amplitude and frequency were pseudo-randomly determined. A portion of the target trajectory is shown by small rectangles in Figure 3.

The relationship between cursor position and the joystick angle was

\[
\begin{pmatrix}
  x \\
  y 
\end{pmatrix} = 0.12 \begin{pmatrix}
  \cos \theta & \sin \theta \\
  -\sin \theta & \cos \theta 
\end{pmatrix} \begin{pmatrix}
  p \\
  q 
\end{pmatrix}
\]

where \((x, y)\) denotes the cursor’s screen coordinates (visual angle: \(^{\circ}\)) and \((p, q)\) denotes the deviational angles of the joystick from the vertical axis in the left-right and the back-forth directions. \(\theta\) was a rotational angle of 0°, 60°, 110°, or 160°. Figure 1 illustrates the correspondence between the direction of joystick movement and that of cursor movement under each rotation.

**Procedures**

All subjects underwent the following: 1) five days of long-term training for the 60° and the 160° joysticks; 2) a pre-test; 3) 25 min of short-term training for the 110° joystick; and 4) a post-test. Brain activity was scanned in the pre- and post-tests.

**Five Days of Long-Term Training for the 60° and the 160° Joysticks**

The subjects were trained for the tracking task under 60° and 160° conditions. During the five-day training sessions, brain activity was not scanned but the subjects performed the tasks lying on a bed just as they would later do in the magnetic resonance imaging (MRI) scanner. Within each session, a tracking task period (1.6 min) and a rest period (18 sec) were alternately repeated three times without changing the joystick type. As shown in Figure 2a, the subjects used the 0° joystick in the first session each morning and were then trained for either the 60° or 160° joystick in the subsequent eight sessions. Breaks between sessions lasted 5-10 min. After a 1-2 hour lunch break, the subjects used the 0° joystick in the first session of each afternoon, and were subsequently trained for the other 60° or 160° joystick in eight sessions. The accumulated tracking time over the five days was 192 min for the 60° and 160° joysticks.
Fig. 1 – Relationship between the direction of joystick movements and that of cursor movements on the screen under each rotation. Black arrows indicate the direction in which subjects moved the joystick, while white arrows indicate the direction of the resulting cursor movement. Numbers indicate correspondence between the black and the white arrows.

Fig. 2 – A graphic representation of the procedures in the present experiment.
Pre-Test

In a pre-test, the subjects used the 0°, 60°, 110°, and 160° joysticks in one session. The order of the tracking periods using these joysticks was counterbalanced between the sessions. An example of the session was three repetitions of [rest → 60° → 0° → 110° → 0° → 160° → 0°] as illustrated in Figure 2b. The rest period lasted 24 sec, while the other periods lasted 48 sec. When the subjects used the 0° joystick, the target velocity was adjusted so that the tracking error was equalized to the error for the joystick used in the preceding period (see Equalization of Tracking Error). The subjects underwent four sessions.

25 Minutes of Short-Term Training for 110° Joystick

One session (7.2 min) was comprised of two repetitions of [rest → 110° → 0° → 110° → 0°] as shown in Figure 2c. The rest periods lasted 24 sec, while each tracking period lasted 48 sec. The subjects underwent eight sessions with 5-10-minute breaks between sessions. The accumulated tracking time for 110° joystick was short (25 min) in
comparison to that for 60° or 160° in the five days of long-term training (192 min).

Post-Test

The subjects underwent the same sessions as in the pre-test.

Analysis of Behavioral Data

The cursor, the target, and the joystick angles were sampled at 1 kHz. The distance between the cursor and the target at each sampling point \((\sqrt{(x_t - x_c)^2 + (y_t - y_c)^2})\) was accumulated over 5.3 sec (position tracking error). A velocity tracking error \((\sqrt{(x_t - x_v)^2 + (y_t - y_v)^2})\) was also accumulated over 5.3 sec. Because the results obtained from the velocity error were similar to those from position tracking error, we only present the position tracking error in this article.

Equalization of Tracking Error

In our previous study (Imamizu et al., 2000), two types of activities related to learning were observed in the cerebellum. One reflected error signals that guide the learning acquisition of internal models and the other reflected an acquired internal model. The former activity was so strongly distributed over the cerebellum that it also blurred the latter activity. Following the procedure used in the previous study, tracking error during manipulation of the normal (0°) joystick was equalized to that during the rotated joystick using a linear relationship between tracking error and target velocity. This procedure allowed us to distinguish internal model activity for the novel tools from internal models and the other reflected an acquired activity reflecting the error signal.

Before functional imaging, the subjects performed the tracking task using the 0° joystick at various target velocities \((V_0)\) that ranged from 0.0427 to 0.0783°/second at 0.0021 intervals. A linear relationship between \(V_0\) and the tracking error \((E)\) was derived by a least squares method for each subject:

\[
E = aV_0 + b.
\]

In the pre- and post-tests, the target velocity was adjusted using the estimated values \(a\) and \(b\) in the 0° baseline condition, so that the error was equal to the mean error \((\bar{E})\) in the preceding 60°, 110°, or 160° test periods:

\[
V_0 = \frac{\bar{E} - b}{a}
\]

The averaged target velocity was .0652°/second in the pre-test and .0543°/second in the post-test.

MRI Acquisition

A 1.5 T MRI scanner (Shimadzu-Marconi) was used to obtain blood oxygen level dependent (BOLD) contrast functional images. Images weighted with the apparent transverse relaxation time were obtained with an echo-planar imaging sequence (repetition time = 5.3 sec, echo time = 65 msec, flip angle = 90°). A total of 158 sequential whole brain volumes \((64 \times 64 \times 44\) voxels at 3.4-mm isotropic resolution) were acquired in each session. High resolution anatomical images of all subjects were also acquired with a T1 weighted sequence.

MRI Analysis

We used Statistical Parametric Mapping 99 (SPM 99) software (http://www.fil.ion.ucl.ac.uk/spm/) for image processing and analysis. The first two volumes of images were discarded to allow for T1 equilibration while the remaining 156 image volumes were realigned to a reference volume. The first volume was chosen as the reference to minimize the difference between the realigned functional images and the anatomical image that was acquired immediately after subjects entered the MRI scanner. The realigned images were normalized to the Montreal Neurological Institute (MNI) reference brain. The data were spatially smoothed with a Gaussian kernel with a 7 mm full-width half-maximum (FWHM). The voxel time series were temporally smoothed with a Gaussian filter (FWHM of 4 sec).

We conducted a multiple regression analysis to find regions related to the manipulation of the three types of joysticks.

\[
v_i^k = \alpha w_i^k + \beta x_i^k + \gamma y_i^k + \delta z_i^k + e_i
\]

Here, \(v_i^k\) denotes the functional magnetic resonance imaging (fMRI) signal at the \(i\)-th voxel in the \(k\)-th scan, and \(w, x, y\) and \(z\) are explanatory variables corresponding to the use of the 0° (error equalized), 60°, 110°, and 160° joysticks, respectively. They were assigned 1 if the scan corresponded to their joystick type and 0 otherwise.

We performed a random effect analysis comprising two stages. In the first stage analysis, parameters were estimated for the differences of interest \((\beta - \alpha, \gamma - \alpha\) and \(\delta - \alpha\)) for each subject. They were then entered into a second-level analysis to test whether parameters for the 60°, 110°, or 160° joysticks were significantly larger than parameters for the error-equalized 0° joystick \((\beta - \alpha > 0, \gamma - \alpha > 0\) and \(\delta - \alpha > 0)\), using a one-sample \(t\)-test across subjects \([t(13) > 3.85, p < .05\) uncorrected\].

Quantitative Analysis of Activity Change from Pre- to Post-Test

We searched for voxels in which activation was significantly associated with all three types of joystick. Then we identified the anatomical volume of interest (VOI) including the voxel according to (Tzourio-Mazoyer et al., 2002) and investigated the change in activated volume separately for each
VOI. The voxels were found in the premotor cortex and regions near the IPS. The premotor VOI was defined as Broadmann area 6, excluding the medial cortex but including both the ventral (PMv) and dorsal (PMd) parts. The parietal VOI comprised the superior parietal gyrus, the angular gyrus, the supramarginal gyrus, and the precuneus. The cerebellum was divided into upper and lower parts by a horizontal plane \((z = -40)\) that roughly corresponded to the horizontal fissure.

**RESULTS**

**Behavioral Data**

Figure 3 shows a representative target and cursor trajectory from a subject’s data while using each of the rotated angles. Although the target trajectories were randomly determined and thus different for the four angles, the relationship between the target motion and the corresponding cursor motion was qualitatively the same.

Figure 4a shows how the tracking errors changed during long-term training for the 60° (red) and 160° (blue) joysticks before scanning. The errors for both joysticks decreased as the accumulated tracking time increased. A repeated measures analysis of variance (ANOVA) for errors indicated a significant time effect for the 60° joystick \([F (1, 39) = 1.76, p < .005]\) and for the 160° joystick \([F (1, 39) = 3.25, p < .0001]\), suggesting that learning had occurred.

Figure 4b shows errors during the pre-test, the short-term training for the 110° joystick, and the post-test. Using a paired \(t\)-test, we examined whether there was a significant difference between the errors in the pre- and post-tests. No significant
difference was observed for the 60° or 160° joystick [60°: t(13) = .17, 160°: t(13) = 1.57] but a significant decrease was observed for the 110° joystick [t(13) = 3.83, p < .001]. Thus, a significant effect of short-term training for the 110° training could be identified.

In the pre- and post-tests, target velocity was adjusted during the use of the 0° joystick to equalize tracking error to the error during use of the 60°, 110° and 160° joysticks (see Equalization of Tracking Error in Methods). There was no significant difference between these two equalized errors during both the pre-test and post-test for each type of joystick [60°: t(13) = .64, 110°: t(13) = 1.19, 160°: t(13) = .97 in the pre-test; 60°: t(13) = .24, 110°: t(13) = 1.55, 160°: t(13) = .57 in the post-test], suggesting that error-equalization succeeded. The activations shown below were obtained by subtracting the corresponding 0° activation from the 60°, 110°, or 160° activation as explained in the MRI analysis section.

**fMRI Data**

**Change of Three Types of Activation from Pre- to Post-Test**

Figures 5a and 5b show the 60° (red), 110° (green), and 160° (blue) activations in the pre-test [t(13) > 3.85, p < .05 uncorrected, cluster size > 10 voxels]. Activations are shown in left-superior-posterior view (a and c) and in right-superior-anterior view (b and d). The cerebrum and the cerebellum are separated into the upper and lower parts of each panel.
In the post-test, the red (60°) regions decreased in various regions, especially in the left and right intraparietal regions and the cerebellum. The green (110°) regions decreased in the cerebral cortex but increased in the inferior parts of the cerebellum. In the right ventral premotor cortex, in the pre-test the dominant color was blue (160°) but it was green (110°) in the post-test.

We searched for regions where the three types of activation (for 60°, 110° and 160°) overlapped in the pre-test and found them in the left premotor region, the left and right intraparietal regions, and the superior left and inferior right cerebellum. The Reorganization of internal models

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<td>Supramarginal gyrus</td>
<td>(40) R (40, – 36, 42)</td>
</tr>
<tr>
<td>Supramarginal gyrus</td>
<td>(40) L (– 32, 38)</td>
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<td>Limbic lobe</td>
<td>Median cingulate and paracingulate gyri</td>
<td>(6) R (8, – 6, 48)</td>
</tr>
<tr>
<td>Insula lobe</td>
<td>Insula</td>
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</tr>
<tr>
<td>Subcortical gray nuclei</td>
<td>Thalamus</td>
<td>(42, 14, 0)</td>
</tr>
<tr>
<td>Lenticular nucleus pallidum</td>
<td>L (8, 0, 2)</td>
<td>5.52</td>
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<tr>
<td>Cerebellum</td>
<td>Crus 1</td>
<td>L (– 36, – 46, – 36)</td>
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<tr>
<td>Lobule 3</td>
<td>L (30, – 70, – 30)</td>
<td>5.91</td>
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<td>Lobule 4-5</td>
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<td>Vermis 9</td>
<td>R (10, – 54, – 32)</td>
<td>5.11</td>
</tr>
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lng. 5c and 5d show the activations in the post-test. The red (60°) regions decreased in the post-test in various regions, especially in the left and right intraparietal regions and the cerebellum. The green (110°) regions decreased in the cerebral cortex but increased in the inferior parts of the cerebellum. In the right ventral premotor cortex, in the pre-test the dominant color was blue (160°) but it was green (110°) in the post-test.

We searched for regions where the three types of activation (for 60°, 110° and 160°) overlapped in the pre-test and found them in the left premotor region, the left and right intraparietal regions, and the superior left and inferior right cerebellum. The t-value weighted centers of gravity of those overlapping regions were (– 27, – 5, 51), (– 37, – 42, 45), (38, – 35, 41), (– 23, – 66, – 25) and (14, – 61, – 51), respectively. The number of voxels included in the regions were 6, 4, 19, 12 and 3. These regions are assumed to be closely related to the manipulation of the rotated joysticks.

Change of 110° Activity from Pre- to Post-Test

We searched for regions where the three types of activation (for 60°, 110° and 160°) overlapped in the pre-test and found them in the left premotor region, the left and right intraparietal regions, and the superior left and inferior right cerebellum. The t-value weighted centers of gravity of those overlapping regions were (– 27, – 5, 51), (– 37, – 42, 45), (38, – 35, 41), (– 23, – 66, – 25) and (14, – 61, – 51), respectively. The number of voxels included in the regions were 6, 4, 19, 12 and 3. These regions are assumed to be closely related to the manipulation of the rotated joysticks.

Change of 110° Activity from Pre- to Post-Test

Table I lists the 110° activation peaks and peak t-values. A number of cerebral regions were activated in the pre-test but not all were significantly activated in the post-test. A decrease
of activation was prominent in the prefrontal regions (area 6 in the superior frontal gyrus; areas 8, 45, and 46 in the middle frontal gyrus) and in the parietal regions. In contrast, an increase of activity was observed in the premotor (area 44 in the opercular part of the inferior frontal gyrus and area 6 in the superior frontal gyrus at \( z = 74 \) level) and in the SMA. The total number of voxels did not change drastically in the cerebellum. However, activation shifted from the superior parts (crus 1 and lobules 3-6) to the inferior parts (lobules 7b-9).

Figures 6a-6e show representative slices from the above regions. The panels in Figure 6a show areas 45 and 46 in the prefrontal regions where significant activation was observed in the pre-test but not in the post-test. Figures 6b and 6c include the dorsal and ventral premotor regions, respectively. Activation in the dorsal and ventral premotor regions increased in the post-test. Figure 6d shows the intraparietal regions where the 60° (red) and 110° (green) activations decreased in the post-test. Figure 6e shows the cerebellar activity in
the coronal sections. The 60° activation decreased in the superior parts, and the 110° activation increased in the inferior parts.

Quantitative Analysis of Change in Activity

We compared activated volume in the pre-test to that of the post-test for each subject for anatomical regions including the overlap among the three types of activation (Figure 7). The activated volume was measured in activation map of each subject (p < .001 uncorrected). This analysis was conducted not only in the regions where the overlap was found but also in the homologous region on the other side of the hemisphere. Below, we will describe increases or decreases thresholded at p < .10.

Although marginally significant (not significant in the strict sense) results pass this threshold, we will report them to indicate trends in change of activity.

Regarding the premotor regions, a significant volume decrease was found in the left premotor [t(13) = 1.89, p < .04, paired t-test] for the 60° activation. In the left and right intraparietal regions, a significant or marginally significant decrease was identified for the 60° activation [left: t(13) = 1.45, p < .09; right: t(13) = 1.71, p < .06] and the 110° activation [left: t(13) = 1.85, p < .05; right: t(13) = 1.63, p < .06]. The decrease in the 110° activation in the parietal regions was consistent with Table I. In the superior part of the cerebellum, the volume for the 60° activation decreased in the post-test [left: t(13) = 1.92, p < .04; right: t(13) = 1.54, p <...
DISCUSSION

In summary, the 110° activation in the frontal and parietal regions decreased in the post-test whereas it increased in the premotor (in the precentral gyrus and the opercular part of the inferior frontal gyrus) regions, in the SMA, and in the inferior part of the cerebellum. The 60° activation decreased in most of the brain regions. Regarding 160° activation, no significant change in activity was observed.

As mentioned in the Introduction, we previously investigated cerebellar activity on subjects who learned to use a 120° rotated mouse and identified activity reflecting an internal model separate from activity reflecting performance error. We used the same method to equalize tracking error as in the current experiment (Imamizu et al., 2000). We also previously investigated cerebellar activity after learning the use of two novel tools (a rotated mouse and a velocity mouse whose cursor velocity was proportional to the mouse’s position). The activities of the two different tools were spatially segregated with slight overlap, suggesting that multiple internal models are acquired in a modular fashion (Imamizu et al., 2003).

The first advantage of possessing multiple internal models is the reduction of interference between different learning epochs, which enables the rapid switching of skilled behaviors. The second advantage is being able to cope with an entirely new environment by adaptively mixing pre-existing motor primitives as multiple internal models. However, the central nervous system (CNS) must solve a selection problem: deciding which internal models are appropriate for usage and learning under a given context of sensory-motor behaviors. MOSAIC models have been proposed for the learning and switching of internal models (Wolpert and Kawato, 1998; Kawato and Wolpert, 1998b; Wolpert et al., 1998, 2003; Wolpert and Ghahramani, 2000; Haruno et al., 2001; Doya et al., 2002). In this model, output from multiple internal models are mixed in proportion to their appropriateness for the current context. Therefore, an enormous repertoire of behavior can be generated even if the number of internal models is limited, and many situations that we encounter may be derived from a combination of previously experienced situations. A recent study (Imamizu et al., 2004) suggests that the dorsolateral prefrontal (area 46) and the parietal regions contribute to the selection and combination (‘blending’) of multiple internal models. A PET study on monkeys also found that fronto-cerebellar interaction contributes to switching tools when monkeys use several tools simultaneously (Obayashi et al., 2002). A recent neuroanatomical study (Kelly and Strick, 2003) found loop connections between area 46 and the cerebellum, which is consistent with these results.

Based on a series of previous studies that we have conducted, the results of our current study can be interpreted as follows. Before scanning brain activity, an internal model for the 60° joystick and one for the 160° joystick were acquired in the cerebellum as a result of long-term training. In the pre-test, it was assumed that the CNS copes with the new 110° joystick by blending the output from the 60° and the 160° internal models. Activation in area 46 and the parietal regions contributed to this blending process. An increase in 110° activity was observed in the premotor regions, in the SMA, and in the inferior part of the cerebellum in the post-test in comparison to pre-tests, suggesting that an internal model for the 110° joystick was acquired in these regions. For the following reasons, it is assumed that internal models are acquired in the inferior part of the cerebellum. First, group analysis (random effect analysis) indicated that the volume increase was the largest in lobule 7b of the inferior cerebellum (from 15 to 686 + 11 voxels, Table I). Second, a marginally significant increase of activation was identified only in the inferior cerebellum among the regions where the quantitative analysis was conducted (Figure 7). As mentioned above, a previous study that we conducted indicated that functional connectivity between the lateral parts of the cerebellum and the premotor regions increased after acquisition of the internal model, which suggests that the acquired internal models send outputs to the premotor regions (Tamada et al., 1999). Electrophysiological studies (e.g., Sasaki et al., 1977) and neuroanatomical studies (Middleton and Strick, 1994; Kelly and Strick, 1998; Dun and Strick, 2003) have revealed reciprocal connections between the cerebellar and premotor regions. It is supposed that internal models for the 60° and 160° joysticks had already been acquired in the pre-test and that these models sent outputs to the premotor regions. The connectivity probably further increased in the post-test after acquisition of the 110° internal model. Our study suggested that the SMA also receives outputs from the acquired internal models. The 60° and the 110°
internal models contributed to manipulation of the 110° joystick in the pre-test, but it is unknown whether these models contributed to learning the 110° internal models.

The results of our current experiments do not contradict the following computational interpretations. When encountering a new tool, the CNS attempts to cope with it by blending outputs from internal models previously acquired. The prefrontal and parietal regions receive outputs from the internal models and contribute to the blending. However, after the acquisition of a proper internal model, the output of the internal model is directly sent to the premotor regions. Our findings suggest that the acquisition of a new cognitive function causes reorganization in the lateral cerebellum and changes global information flow in the cerebro-cerebellar communication loop.

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