



Rethinking Motor Learning and Savings in Adaptation Paradigms: Model-Free Memory for Successful Actions Combines with Internal Models

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SUMMARY

Although motor learning is likely to involve multiple processes, phenomena observed in error-based motor learning paradigms tend to be conceptualized in terms of only a single process: adaptation, which occurs through updating an internal model. Here we argue that fundamental phenomena like movement direction biases, savings (faster relearning), and interference do not relate to adaptation but instead are attributable to two additional learning processes that can be characterized as model-free: use-dependent plasticity and operant reinforcement. Although usually "hidden" behind adaptation, we demonstrate, with modified visuomotor rotation paradigms, that these distinct model-based and model-free processes combine to learn an errorbased motor task. (1) Adaptation of an internal model channels movements toward successful error reduction in visual space. (2) Repetition of the newly adapted movement induces directional biases toward the repeated movement. (3) Operant reinforcement through association of the adapted movement with successful error reduction is responsible for savings.

INTRODUCTION

Skilled motor behaviors outside the laboratory setting require the operation of multiple cognitive processes, all of which are likely to improve through learning (Wulf et al., 2010; Yarrow et al., 2009). Several simple laboratory-based tasks have been developed in an attempt to make the complex problem of motor learning more tractable. For example, error-based paradigms have been used extensively to study motor learning in the context of reaching movements (Debicki and Gribble, 2004; Flanagan et al., 2003; Held and Rekosh, 1963; Imamizu et al., 1995; Krakauer et al., 1999; Lackner and Dizio, 1994; Malfait et al., 2005; Miall et al., 2004; Pine et al., 1996; Scheidt et al.,

2001; Shadmehr and Mussa-Ivaldi, 1994). In these paradigms, subjects experience a systematic perturbation, either as a deviation of the visual representation of their movements, or as a deflecting force on the arm, both of which induce reaching errors. Subjects then gradually correct these errors to return behavioral performance to preperturbation levels.

Error reduction in perturbation paradigms is generally thought to occur via adaptation: learning of an internal model that predicts the consequences of outgoing motor commands. When acting in a perturbing environment, the internal model is incrementally updated to reflect the dynamics of the new environment. Improvements in performance are usually assumed to directly reflect improvements in the internal model. This learning process can be mathematically modeled in terms of an iterative update of the parameters of a forward model (a mapping from motor commands to predicted sensory consequences) by gradient descent on the squared prediction error (Thoroughman and Shadmehr, 2000), which also can be interpreted as iterative Bayesian estimation of the movement dynamics (Korenberg and Ghahramani, 2002). This basic learning rule can be combined with the notion that what is learned in one direction partially generalizes to neighboring movement directions (Gandolfo et al., 1996; Pine et al., 1996), leading to the so-called state space model (SSM) of motor adaptation (Donchin et al., 2003; Thoroughman and Shadmehr, 2000). Despite its apparent simplicity, the SSM framework fits trial-to-trial perturbation data extremely well (Ethier et al., 2008; Huang and Shadmehr, 2007; Scheidt et al., 2001; Smith et al., 2006; Tanaka et al., 2009). In addition, parameter estimates from state-space model fits also predict many effects that occur after initial adaptation such as retention (Joiner and Smith, 2008) and anterograde interference (Sing and

The success of the SSM framework has led to the prevailing view that the brain solves the control problem in a fundamentally model-based way: in the face of a perturbation, control is recovered by updating an appropriate internal model, which is then used to guide movement. An alternative view is that a new control policy might be learned directly through trial and error until successful motor commands are found. No explicit model of the perturbation is necessary in this approach and thus it



can be described as model-free. This distinction between model-free and model-based learning originates from the theory of reinforcement learning (Kaelbling et al., 1996; Sutton and Barto, 1998). However, the dichotomy is applicable in any scenario where a control policy must be learned from experience, not just when explicit rewards are given. If learning in perturbation paradigms were purely model-free, one would expect substantial trial-to-trial variability in movements. However, such exploratory behavior is not usually observed; in fact, it is only seen if subjects receive nothing but binary feedback about success or failure of their movements (Izawa and Shadmehr, 2011).

Despite the success of SSMs in explaining initial reduction of errors, there are phenomena in adaptation tasks that these models have difficulty accounting for. In particular, relearning of a given perturbation for a second time is faster than initial learning, a phenomenon known as savings (Ebbinghaus, 1913; Kojima et al., 2004; Krakauer et al., 2005; Smith et al., 2006; Zarahn et al., 2008), whereas a basic single-timescale SSM predicts that learning should always occur at the same rate, regardless of past experience (Zarahn et al., 2008). Although SSM variants that include multiple timescales of learning (Kording et al., 2007; Smith et al., 2006) are able to explain savings over short timescales, this approach fails to predict the fact that savings still occurs following a prolonged period of washout of initial learning (Krakauer et al., 2005; Zarahn et al., 2008). Beyond SSMs, there are other potential ways to explain savings and still remain within the framework of internal models. For example, more complex neural network formulations of internal model learning can exhibit savings despite extensive washout (Ajemian et al., 2010), owing to redundancies in how a particular internal model can be represented. Another possible explanation is that rather than updating a single internal model, savings could occur by concurrent learning and switching between multiple internal models, with apparent faster relearning occurring because of a switch to a previously learned model (Haruno et al., 2001; Lee and Schweighofer, 2009). The core idea in all of these models is that savings is the result of either fast reacquisition or re-expression of a previously learned internal model; i.e., they all explain savings within a model-based learning framework.

An entirely different idea is that savings does not emerge from internal model acquisition but instead is attributable to a qualitatively different form of learning that operates independently. We hypothesize that savings reflects the recall of a motor memory formed through a model-free learning process that occurs via reinforcement of those actions that lead to success, regardless of the state of the internal model. This idea is consistent with the suggestion that the brain recruits multiple anatomically and computationally distinct learning processes that combine to accomplish a task goal (Doya, 1999). We posit an operant process whereby the movement on which adaptation converges in hand space is reinforced because it is associated with successful target attainment in the context of a perturbation; this operant memory should not be affected by washout of an internal model. In this formulation, savings would result from accelerated recall of the reinforced action rather than of an internal model.

Support for the idea that a memory for actions exists independently of internal models comes from experiments in which repetition of a particular action leads future movements to be biased toward that action (Classen et al., 1998; Jax and Rosenbaum, 2007; Verstynen and Sabes, 2011). Since these experiments do not entail any change in the dynamics of the environment, these biases cannot be explained in terms of the framework of internal models. Instead, they reflect a form of model-free motor learning. More recently it has been shown that biases can be observed in parallel with acquisition of an internal model along the task-irrelevant dimension in a redundant task (Diedrichsen et al., 2010). The term that has been used for these repetition-induced biases is use-dependent plasticity (Bütefisch et al., 2000; Classen et al., 1998; Diedrichsen et al., 2010; Krutky and Perreault, 2007; Ziemann et al., 2001). Here we will argue that the process underlying savings is also model-free but distinct from use-dependent plasticity.

We hypothesized that multiple learning processes can combine along the task-relevant dimension of an adaptation task. We sought to dissociate model-based (adaptation) and model-free (use-dependent plasticity and operant reinforcement) learning processes using variants of a visuomotor rotation paradigm that either eliminated or exaggerated movement repetition in the setting of adaptation. Our prediction was that, following adaptation in the absence of repetition, model-free learning processes would not be engaged and subjects would exhibit neither savings nor biases in execution of subsequent movements. Conversely, we predicted that both savings and movement biases would be more prominent when repetition is exaggerated in the context of error reduction.

RESULTS

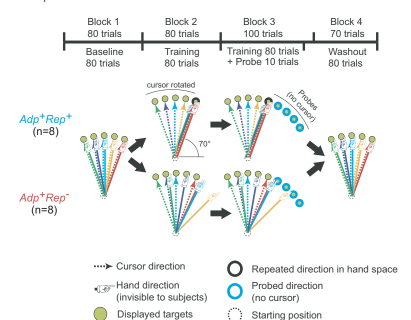
Experiment 1: Movement Repetition Caused by Adaptation Induced Large Directional Biases

We first sought to test the hypothesis that biases can be induced along the task-relevant dimension (movement direction) of a visuomotor rotation task in the setting of adaptation (Figure 1A). We compared two groups of subjects that were exposed to identical, uniform distributions of counterclockwise ("+") visuomotor rotations (mean = $+20^{\circ}$, range = $[0^{\circ}, +40^{\circ}]$) (see Figure S1B available online). The protocol for the first group was predicated on the idea that adaptation itself, by converging on a single movement direction that is then repeated, can induce directional biases. We wished to exaggerate this purported asymptotic process in order to unmask it by designing an adaptation protocol for which the adapted solution in hand space would be the same for all visual target directions (Figure 1A). Specifically we introduced a target-dependent structure to the sequence of rotations such that the ideal movement in hand space was always in the 70° direction. In other words, cursor feedback of a movement made toward a target at θ was rotated by $+(\theta - 70)^{\circ}$ (Figure 1B). We named this group $Adp^{+}Rep^{+}$ and refer to the 70° movement direction in hand space as the "repeated direction" (Figure 1A). It should be noted that although adaptation is not a prerequisite for biases to occur (Diedrichsen et al., 2010; Verstynen and Sabes, 2011), here the idea was to

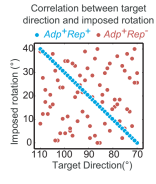


Α

Experiment 1 Protocols



В



exploit adaptation to induce repetition of a particular movement

In the second group, Adp^+Rep^- (i.e., adaptation-only), which served as a control, we sought to induce pure adaptation without the possibility of repetition-induced biases, which was accomplished by sampling from the same perturbation distribution and randomly varying the rotations at each target so that the solution in hand space was never repeated for any given target (Figures 1A and 1B). Subjects in Adp⁺Rep⁻ were expected to counterrotate by -20° on average (Scheidt et al., 2001), making 70° movements in hand space on average for all visual targets as the result of adaptation alone.

The imposed rotations resulted in reaching errors that drove both Adp^+Rep^- and Adp^+Rep^+ to adapt (Figures 2A and 2B).

Figure 1. Protocols for Experiment 1

(A) Adapted movement directions in hand space are represented by solid "pointing hand" arrows, corresponding cursor movement directions in visual space are represented by dotted arrows in the same color. For Adp+Reptraining, cursor feedback was rotated by random, counterclockwise angles sampled from a uniform distribution ranging from 0° to +40°. For Adp+Rep+ training, cursor feedback was rotated by a target-specific angle, sampled from the same uniform distribution as Adp+Rep-, such that the hand always had to move in the 70° direction for the cursor to hit the target (repeated direction in hand space). In probe trials subjects had to move to targets shown clockwise from the training targets without cursor feedback. Numbers and locations of targets are schematic and not to scale.

(B) In Adp+Rep-, the imposed rotation was randomly selected every time the subject visited each target. In Adp+Rep+, the rotations were structured so that the adapted hand movement was always toward the 70° direction in hand space.

State-space models have been used extensively in adaptation studies and have shown good fits to trial-to-trial data (Donchin et al., 2003; Huang and Shadmehr, 2007; Scheidt et al., 2001; Smith et al., 2006; Tanaka et al., 2009; Thoroughman and Shadmehr, 2000). We reasoned that if we had succeeded in creating a condition that only allowed adaptation, Adp+Rep-, then a statespace model that describes the process of internal model acquisition would simulate the empirical data well. In contrast, in Adp+Rep+, we predicted that we would obtain a good state-space model fit during initial leaning but that subsequently subjects' performance would be better than predicted because of the presence of additional model-free learning processes that become engaged through repetition of the same movement. We obtained rotation learning parameters and the directional generalization function width from our previously published data (Tanaka et al., 2009) and used these to generate simulated hand directions for the target sequences presented in Adp+Rep+ and

Adp⁺Rep⁻ during training (Figures 2C and 2D, "adapt-only sim"). The state-space model was an excellent predictor of the empirical data for Adp^+Rep^- (r² = 0.968, Figure 2C), which supports our assumption that asymptotic performance in Adp⁺Rep⁻ can be completely accounted for by error-based learning of an internal model alone; subjects rotated their hand movement by an average of $-13.97 \pm 1.41^{\circ}$ (mean \pm SD) (the vertical displacement from the naive line in Figure 2C), or about 70% adaptation on average for all targets.

For Adp⁺Rep⁺, the adaptation model was able to predict hand directions relatively well in the early phase of training ($r^2 = 0.753$) but then began to fail as subjects developed a directional bias beyond what was expected from adaptation alone $(r^2 = -0.502)$ (Figure 2D, asymptotic training). This suggests



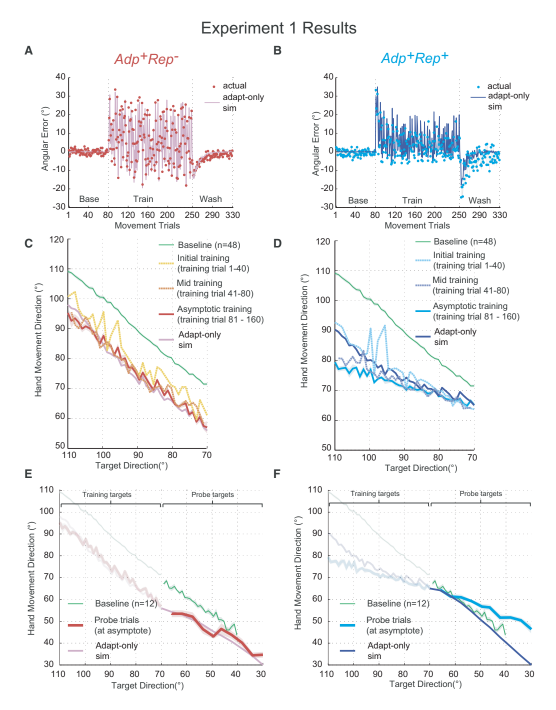


Figure 2. Group Average Results for Experiment 1

Left: Adp⁺Rep⁻; right: Adp⁺Rep⁺.

(A and B) Time courses for empirical trial-to-trial data (dots) and adaptation (state-space) model simulations (lines). Errors were computed as the angular separation between cursor and target direction (shadings indicate SEM).

(C and D) Hand-movement direction versus displayed target direction for the initial, middle, and asymptotic phases of training: both data and simulation at asymptote shown. For comparison, baseline data from all subjects are also plotted (green line). Shading indicates SEM. The two "peaks" in initial training line show that the performance of the first two trials in training was close to naive performance.

(E and F) Hand-movement direction versus target direction for generalization probes: both data and simulation shown. Panels from (C) and (D) are replotted in faded colors. Baseline performance to the probe target directions for a separate group of subjects is plotted (green) for comparison. Shading indicates SEM.



that errors were first reduced through adaptation but then were further reduced through mechanisms other than adaptation.

The divergence between the data and the model in Adp^+Rep^+ had a particular structure: a bias toward the repeated direction. Indeed, at training asymptote, movement directions in hand space for Adp^+Rep^+ were more tightly distributed around the repeated direction (mean SD = $4.9 \pm 0.4^\circ$, mean \pm SEM) when compared to Adp^+Rep^- (mean SD = $11.7 \pm 0.45^\circ$, t(14) = -11.95, p < 0.001). This tight distribution of hand movements at asymptote constituted our key step for induction of use-dependent learning (distribution shown in Figure S1D), which we posited would manifest as a movement bias toward the mean of the hand movement distribution at the end of training (i.e., toward the repeated direction). The mean movement direction at the end of training across subjects was $76.0 \pm 2.1^\circ$ (mean \pm SD) for Adp^+Rep^- (Figure S1D) and the mean movement direction at the end of training was $71.6 \pm 1.3^\circ$ (mean \pm SD) for Adp^+Rep^+ .

We tested for generalization in a mirror subset of untrained probe targets arrayed evenly and clockwise of the repeated direction (Figure 1A, Block 3). No cursor feedback was provided in these trials. Our previous work has demonstrated that generalization for adaptation alone falls off as a function of angular separation away from the training direction (Donchin et al., 2003; Gandolfo et al., 1996; Krakauer et al., 2000; Pine et al., 1996; Tanaka et al., 2009); subjects return to their default 0° mapping once they are 45° from the training direction. Within this range, the direction of movements in hand space should always be opposite to the rotation in visual space. In other words, since all the imposed rotations were counterclockwise, all movements toward the probes in hand space should rotate clockwise relative to the target direction. As expected for generalization of adaptation, hand directions in Adp⁺Rep⁻ were clockwise and gradually converged to naive performance and this was predicted well by the state-space model (Figure 2E). However, if we were correct in surmising that the Adp+Rep+ protocol induced biased movements toward the repeated direction then this would predict a similar pattern of directional biases at the probe targets. Adp+Rep+ crossed and began to show an increasing bias away from naive directions as the probe directions moved further away from the repeated direction in hand space (Figure 2F); this is the opposite of the expectation for adaptation but entirely consistent with a bias toward the repeated direction (Verstynen and Sabes, 2011).

Interestingly, the bias generated during Adp^+Rep^+ , which can be plotted as the dependent relationship between displayed targets and hand movement direction, was also apparent during learning, with a slope of 0.32 (\pm 0.03) for the trained targets that was comparable to the slope for the probe targets (0.42 \pm 0.04). To summarize Experiment 1, adaptation to a target sequence that led to movements distributed around the repeated direction in hand space led to a bias toward the repeated direction that was comparable for trained and untrained targets, with increasing absolute size of bias for farther away targets in both directions. These results are opposite of what would be predicted if the observed behavior were solely due to adaptation of an internal model and show that a model-free process based on repeated actions is in operation in Adp^+Rep^+ but not Adp^+Rep^- .

Experiment 2: Savings Occurred following Adaptation-Induced Repetition, but Not with Either Adaptation or Repetition Alone

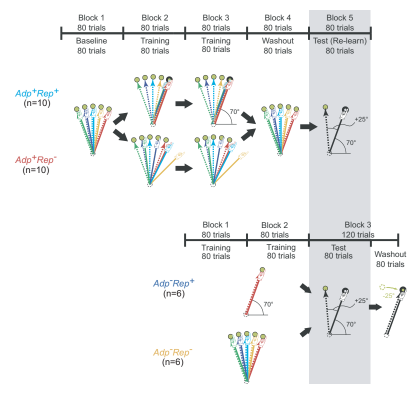
The results of Experiment 1, which showed directional biases in the Adp+Rep+ group, suggested a possible mechanism for savings: subjects in Adp+Rep+ learned to associate the repeated 70° direction movement in hand space with successful adaptation to all targets, i.e., a particular movement in hand space was associated with successful cancellation of errors in the setting of a directional perturbation at all targets. This led us to hypothesize that savings may, at least in part, be attributable to recall of the movement direction that was reinforced at or near asymptote during initial adaptation. The idea is that as readaptation proceeds it will bring subjects within the vicinity of the movement direction that they have previously experienced and associated with successful adaptation; they will therefore retrieve this direction before adaptation alone would be expected to converge on it. Therefore, the prediction would be that postwashout re-exposure to a rotation at a single target would lead to savings for Adp+Rep+ when the readapted solution in hand space is the previously repeated direction, but there would be no savings for Adp^+Rep^- . Also no savings would be predicted after repetition alone (Adp-Rep+) because it would not be associated with (previously successful) adaptation. Finally, a naive group practiced movements in all directions in the absence of a rotation (Adp-Rep-); this group had no error to adapt to and movements to multiple directions would prevent repetition-related directional biases. Thus, Adp⁻Rep⁻ served as a control for the other three groups.

We therefore studied four new groups of subjects who each underwent one of four different kinds of initial training $(Adp^+Rep^+, Adp^+Rep^-, Adp^-Rep^+, Adp^-Rep^-)$. The two Adp^+ groups had a washout block after training and all four groups were tested with a $+25^{\circ}$ rotation at the 95° target (Figure 3). That is, the movement solution in hand space for the test session was again the 70° direction. We chose a $+25^{\circ}$ rather than a $+20^{\circ}$ rotation in order to increase the dynamic range available to demonstrate savings and because reinforcement should be rotation angle invariant as it is the adaptation-guided direction in hand space that matters. We fit a single exponential function to each subject's data to estimate the rate of error-reduction, expressed as the inverse of the time constant (in units of trial $^{-1}$). Savings would be indicated by a faster error-reduction rate for relearning when compared to naive learning.

We first tested for savings in Adp^+Rep^+ and Adp^+Rep^- . On the first test trial after washout, both Adp^+Rep^+ and Adp^+Rep^- , produced errors close to 25°, which indicated that washout was complete (Adp^+Rep^+ : 23.73 \pm 1.18° (mean \pm SEM); Adp^+Rep^- , 24.20 \pm 2.37, t(18) = -0.340, p = 0.738) (Figure 4A). We fit a single exponential function to each subject's data to estimate the rate of error reduction (Figure 4C). In support of our hypothesis, Adp^+Rep^+ showed significant savings (0.49 \pm 0.08 trial⁻¹, mean \pm sem) when compared to the naive training group Adp^-Rep^- (0.13 \pm 0.02 trial⁻¹) (two-tailed t test, t(14) = 3.495, p = 0.004). In contrast, Adp^+Rep^- (0.12 \pm 0.02 trial⁻¹) were no faster than the naive training control and showed no savings (t(14) = -0.39, p = 0.70) (Figures 4A and 4C). An alternative analysis using repeated-measure ANOVA yielded the same result



Experiment 2 Protocols



(not shown). Indeed, Adp^+Rep^+ had a faster rate of relearning rate than Adp^+Rep^- , (t(18) = 4.62, p < 0.001). We had power of 0.8 (see Experimental Procedures) and thus the negative results are likely true negatives. The effect size we saw for savings is comparable to that in previous studies conducted in our and other laboratories. The time constants are similar to our previous report of savings (Zarahn et al., 2008). While savings is defined as faster relearning rate, it has been measured in various ways in published studies; therefore, we converted reported values in the literature to a percentage increase (i.e., [amount of error reduced in relearning - amount of error reduced in naive] /amount of error reduced in naive). The degree of savings reported in the literature is quite variable. For example, we have previously reported a 20% increase for a 30° visuomotor rotation (Krakauer et al., 2005). For force field adaptation, an estimated 23% increase has been reported (Arce et al., 2010). In Experiment 2, we found a 35% increase in the average amount of error reduced in Adp+Rep+ over the first 20 trials when compared to naive $(Adp^{-}Rep^{-})$ (two-tailed, t(14) = -4.175, p = 0.001). Thus, we saw a marked savings effect for a +25° rotation for Adp⁺Rep⁺, but no savings at all for Adp⁺Rep⁻. This suggests that adaptation alone is insufficient to induce savings.

There are, however, two potential concerns with the interpretation of Experiment 2. First, the difference between Adp+ Rep⁺ and Adp⁺Rep⁻ might be attributable to the fact that subjects in these two groups might not have adapted to exactly

Figure 3. Adp+Rep-, Adp+Rep+, Adp-Rep+, and Adp Rep Protocols for Experiment 2, Illustrated in the Same Format as Figure 1A

The Adp+ groups initially trained with counterclockwise rotations drawn from a uniform distribution ranged from 0° to +40°. The Adp groups initially trained without rotation. The test block is shown with a gray box background: in all four groups, every subject was tested at the 95° target with a counterclockwise 25° rotation.

the same degree to the 95° target direction during initial training, although the difference was small (approximately 6°). Second, subjects in Adp+Rep were exposed to a 20° rotation but were then tested on 25°, i.e., a larger angle than they adapted to on average, although it has been shown that adaptation to smaller rotation facilitates subsequent adaptation to a larger rotation (Abeele and Bock, 2001). Therefore, we also tested for savings in two additional groups with a +20° rotation in the 90° direction, where the two groups showed comparable degrees of initial adaptation (Adp^+Rep^- : 17.3 ± 0.8°, Adp^+ Rep^+ : 18.51 ± 0.9°, t(14) = -1.047, p = 0.31) (Figures S1A and S1E). Again, Adp+Rep+ had a significantly greater savings than the Adp+ Rep^- (0.15 ± 0.01 trial⁻¹ versus 0.08 ± 0.02 $trial^{-1}$, t(14) = 3.06, p = 0.009) (Figure S1F).

In contrast, no savings was observed for the repetition-only group, $Adp^{-}Rep^{+}$ (Figure 4B); indeed the learning rate was not significantly

different from naive training in $Adp^{-}Rep^{-}$ (0.16 ± 0.04 trial⁻¹ vs. $0.13 \pm 0.02 \text{ trial}^{-1}$, two-tailed t test, t(10) = 0.594, p = 0.565) (Figure 4C). Of note, there was a small bias at the beginning of the test session for Adp⁻Rep⁺, which suggests the development of usedependent plasticity as the result of single direction training; the imposed rotation was 25° but they started with an initial error of 20.54 ± 2.23° (mean ± SEM) whereas the naive control group started at the expected value of $25.36 \pm 1.93^{\circ}$.

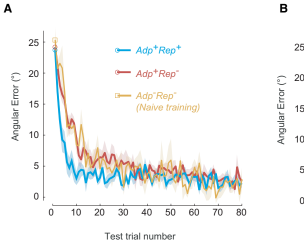
To summarize Experiment 2, an adaptation protocol with movement repetition led to clear savings, whereas neither adaptation alone nor repetition alone led to any savings. These results suggest that the association of movement repetition with successful adaptation is necessary and sufficient for savings.

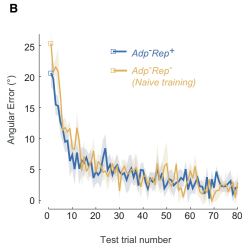
Experiment 3: Savings Occurred for Oppositely Signed Rotations When They Shared the Same Hand-Space Solution

The results of Experiment 2 support the idea that savings is dependent on recall of a repeated solution in hand space. Experiment 2 was designed to exaggerate the presence of model-free reinforcement learning, a process that we argue is present even when the solution in hand space does not map onto multiple directions in visual space. To show that reinforcement also occurs in the more common scenario of one hand-space solution for one visual target, we took advantage of the observation that when rotations of opposite sign are learned sequentially



Experiment 2 Results





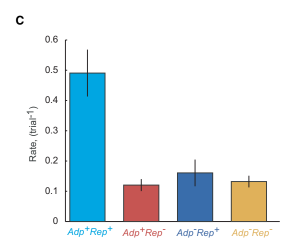


Figure 4. Group Average Results for Experiment 2

(A) Test block learning curves for Adp⁺Rep⁻, Adp⁺Rep⁻, and, Adp⁻Rep⁻. Square and circular markers show the average errors for the first test trial. Errors were computed as the angular separation between cursor and target direction.

(B) Test block learning curves for Adp^-Rep^+ and Adp^-Rep^- . Square markers show the average errors for the first test trial. Errors were computed as the angular separation between cursor and target direction. SEM was omitted for clarity.

(C) Estimated error reduction rates for all four groups during the test block (means of the time constant of a single exponential fit to individual subject data). Error bars indicate SEM.

using the popular A-B-A paradigm (where A and B designate opposite rotations in sign) there is no transfer of savings between A and B, nor subsequent savings when A is relearned (Bock et al., 2001; Brashers-Krug et al., 1996; Krakauer et al., 1999, 2005; Tong et al., 2002; Wigmore et al., 2002). A surprising prediction of our reinforcement hypothesis is that savings should be seen for B after A if the required hand direction is the same for both A and B, even if the two rotations are opposite in sign and learning effects of A are washed out by a intervening block of baseline trials before exposing subjects to B. In this framework, interference (or no savings) in the A-B-A paradigm is attributable to a conflict between the hand-space solutions associated with success for the A and B rotations and not because A and B are opposite in sign in visual space.

Two groups were studied to test the prediction that savings would be seen for a counterrotation after learning a rotation if they shared the same solution in hand space (SAME-SOLN_{hand}) but not if they only shared the same solution in visual space (SAME-SOLN_{visual}) (Figure 5). The SAME-SOLN_{hand} subjects first trained in one target direction (100° target) with a +30° rotation and then, after a washout block, tested in another target direction (40° target) with a counterrotation of -30° . The two different target directions were chosen so that the adapted solution to the two oppositely signed rotations would be the same direction in



Experiment 3 Protocols

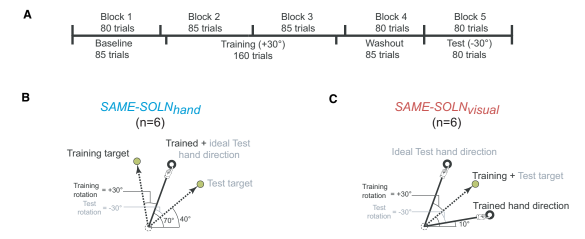


Figure 5. Protocol for Experiment 3

(A) $SAME-SOLN_{hand}$ and $SAME-SOLN_{visual}$ were first trained on a $+30^{\circ}$ rotation then tested on a -30° rotation.

(B) Illustrations of ideal solution in hand space and in visual (cursor) space for $SAME-SOLN_{hand}$. The adapted movement in hand space was the same for both the $+30^{\circ}$ and -30° rotations. Black labels indicate the imposed rotation, the displayed target, and the adapted hand movement direction for initial training with the $+30^{\circ}$ rotation. Gray labels indicate the imposed rotation, the displayed target, and the ideal hand movement direction for the -30° rotation.

(C) Illustrations of ideal solution in hand space and in visual (cursor) space for SAME-SOLN_{visual}. The adapted movement in visual (cursor) space was the same for both the +30° and -30° rotations.

hand space (70°) and so that target separation was sufficient to minimize generalization effects (Tanaka et al., 2009) (Figure 5B). In the SAME- $SOLN_{visual}$ group, subjects first trained in one target direction (40° target) with a + 30° rotation and then, after a washout block, tested in the same target direction with a - 30° rotation. Thus, in this case, the adapted solution for the two rotations was the same direction in visual space, which led to different adapted solutions in hand space (Figure 5C). Baseline and washouts blocks contained equally spaced targets between the 100° and 40° target directions.

The two groups exhibited similar behaviors during initial training (Figure 6). During initial training on $+30^{\circ}$ rotation, $SAME-SOLN_{hand}$ had a learning rate of 0.11 ± 0.04 trial $^{-1}$ (mean \pm SEM) and $SAME-SOLN_{visual}$ had a rate of 0.12 ± 0.04 trial $^{-1}$ (Figure 6C). Consistent with the prediction of operant reinforcement, $SAME-SOLN_{hand}$ showed savings for the -30° rotation after training on $+30^{\circ}$ (Figure 6A); the relearning rate during test $(0.23\pm0.03~{\rm trial}^{-1})$ was significantly faster than initial learning (Figure 6C) (paired one-tailed t(5) = -2.371, p = 0.03). In contrast, no savings were seen for $SAME-SOLN_{visual}$ which had a relearning rate of $0.11\pm0.02~{\rm trial}^{-1}$ during test (Figure 6B) (paired one-tailed t(5) = 0.238, p = 0.411).

Interestingly, in the first test trial of the -30° rotation, $SAME-SOLN_{hand}$ had an average error that was less than the -30° ($-23.34 \pm 0.88^{\circ}$, one-tailed t(5) = 7.56, p < 0.001) while $SAME-SOLN_{visual}$ had an error not significantly different from -30° (t(5) = -0.2, p = 0.849) (Figure 6B). This is consistent with the bias seen in Experiment 1. In summary, the results of Experiment 3 suggest that savings is attributable a model-free operant memory for actions and not to faster relearning or reexpression of a previously learned internal model.

DISCUSSION

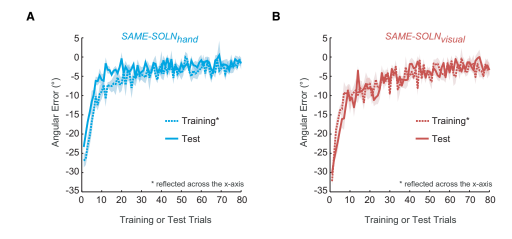
We sought to unmask two model-free learning processes, use-dependent plasticity and operant reinforcement, which we posited go unnoticed in conventional motor adaptation experiments because their behavioral effects are hidden behind adaptation. We found evidence for use-dependent plasticity in the form of a bias toward the repeated direction (i.e., the direction in hand space converged upon by adaptation) for both trained and untrained targets. We found evidence for operant reinforcement in the form of savings: subjects showed faster relearning when rotations of either sign (clockwise or counterclockwise) required an adapted solution that coincided with a previously successful hand movement direction.

Use-Dependent Plasticity

We designed our Adp^+Rep^+ protocol so that adaptation itself would create a narrow distribution of hand movements centered on a particular direction, with the prediction that this would lead to a directional bias via use-dependent plasticity. Adp^+Rep^+ , as expected, did induce a bias toward the mean of the hand movements at asymptote. The hand direction versus target direction relationship was well described with a single linear fit (0 < slope < < 1) that had a close fit to both training and probe targets. These results are consistent with a recent study by Verstynen and Sabes (Verstynen and Sabes, 2011), which showed that repetition alone leads to directional bias. Interestingly, the biases we observed here in the setting of adaptation appear to be larger compared to those induced by repetition alone, which suggests that repetition in the context of reducing errors in response to a perturbation



Experiment 3 Results



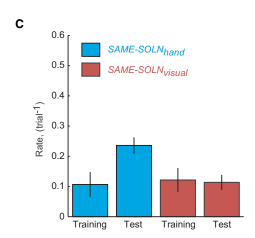


Figure 6. Group Average Results for Experiment 3

(A) Learning curves for initial training and test sessions for SAME-SOLN_{hand}. Shading indicates SEM.

(B) Learning curves for initial training and test sessions for SAME-SOLN_{visual}-

(C) Estimated learning rates for training and test (means of the time constant of a single exponential fit to individual subject data). Error bars indicate SEM.

may in itself generate reward that modulates use-dependent plasticity.

Support for our contention that use-dependent plasticity can be induced during adaptation comes from a force-field adaptation study by Diedrichsen et al. (2010), in which they demonstrated the existence of use-dependent learning in a redundant-task design. In this study, a force channel restricting lateral movements of the hand gradually redirected subjects' hand paths by 8° laterally. However, this had no effect on success in the task because the task-relevant error only related to movement amplitude. Crucially, use-dependent learning occurred in a direction that was orthogonal to the task-relevant dimension, which is why it could be separately identified. Another important result in the study by Diedrichsen and colleagues is that adaptation, pushing in the direction opposite to the channel, occurred in parallel to use-dependent plasticity so that the latter only became apparent after washout of adapta-

tion. The critical difference between our study and that by Diedrichsen and colleagues is that we reasoned that adaptation itself can act like a channel but in the task-relevant dimension; it not only reduces visual error but also guides subjects' hand toward a new path in hand space. Analogous to washing out adaptation in the Diedrichsen et al. study in order to show usedependent plasticity (Diedrichsen et al., 2010), we probed for use-dependent plasticity beyond the range of the expected generalization function for adaptation and found a strong bias toward the repeated direction in hand space.

Savings

Savings is a form of procedural and motor memory that manifests as faster relearning compared with initial learning (Ebbinghaus, 1913; Smith et al., 2006; Zarahn et al., 2008). We reasoned that the reward landscape is not flat during adaptation but rather is increasingly rewarding as the prediction error



decreases; i.e., the hand movement that is induced by the adaptation process will be reinforced through increasing success (decreasing error). Our current results suggest that subjects on second exposure recall the hand direction that was reinforced during the first exposure to the perturbation. Adp^+Rep^+ showed marked savings, whereas Adp^+Rep^- showed no savings, even though they adapted to the same mean rotation. We conclude from Experiment 2 that a reinforcement process was necessary and sufficient for savings, and that use-dependent plasticity is not sufficient for savings.

A set of previously puzzling results reported in visuomotor rotation studies may also be more easily interpreted as arising from an operant model-free mechanism. Savings for a given rotation is disrupted if subjects train with a counterrotation even at prolonged time intervals after initial training and when aftereffects have decayed away (Krakauer et al., 1999, 2005). We propose that persistent interference effects occur because successful cancellation of rotations of opposite sign is associated with different movements in hand space even if the movement of the cursor into the target is the same in visual space. That is, the corresponding motor commands to the same target are distinctly different for oppositely signed rotations. Thus, the association of the same target with different commands in a serial manner, as is done with A-B-A paradigms, could lead to interference as is seen with other forms of paired-associative paradigms. In such paradigms, interference occurs through retrieval inhibition (Adams and Dickinson, 1981; Anderson et al., 2000; MacLeod and Macrae, 2001; Wixted, 2004). Complementary to this explanation for interference, we can predict that there should be facilitation, i.e., savings, for two rotations of opposite sign if they are both associated with the same commands or movements in hand space. This was exactly what we found in Experiment 3: learning a +30° counterclockwise rotation facilitated learning of a -30° clockwise rotation when both rotations required the same directional solution in hand space. This supports the idea that an operant reinforcement process underlies savings and interference effects in adaptation experiments. Furthermore, results from Experiment 3 showed that the directional solution in hand space need not be associated with multiple targets, as in Experiments 1 and 2, for reinforcement to occur; success at a single target, as in Experiment 3 (and in most conventional error-based motor learning paradigms), is sufficient for savings.

Numerous studies suggest that adaptation is dependent on the cerebellum (Martin et al., 1996a, 1996b; Smith and Shadmehr, 2005; Tseng et al., 2007), a structure unaffected in Parkinson's disease (PD), and therefore initial learning in patients with PD would be expected to proceed as in controls, as indeed was recently demonstrated (Bédard and Sanes, 2011; Marinelli et al., 2009). Operant learning is, however, known to be impaired in PD (Avila et al., 2009; Frank et al., 2004; Rushworth et al., 1998; Rutledge et al., 2009; Shohamy et al., 2005). Thus, our contention that initial learning of a rotation occurs through adaptation but savings results from operant learning predicts that patients with PD would show a selective savings deficit in an error-based motor learning paradigm. This is exactly what has been found: patients with PD were able to adapt to initial rotation as well as control subjects but they did not show savings (Bédard and

Sanes, 2011; Marinelli et al., 2009). Thus, our framework of multiple learning processes can explain this otherwise puzzling result. A prediction would be that PD patients would show no difference in learning rates between Adp^+Rep^- and Adp^+Rep^+ protocols, because only adaptation would occur.

Adaptation as Model-Based Learning

Prevailing theories of motor learning in adaptation paradigms have been fundamentally model-based: they posit that the brain maintains an explicit internal model of its environment and/or motor apparatus that is directly used for planning of movements. When faced with a perturbation, this model is updated based on movement errors and execution of subsequent movements reflects this updated model (Shadmehr et al., 2010). We wish to define adaptation as precisely this model-based mechanism for updating a control policy in response to a perturbation. Adaptation does not invariably result in better task performance. For example, in a previous study we showed that adaptation to rotation occurs despite conflicting with explicit task goals (Mazzoni and Krakauer, 2006). In the current study, hyper- or overadaptation occurred to some targets due to unwanted generalization; this was why the steady-state predicted by the state-space model for Adp⁺Rep⁺ showed that subjects adapted past the 70° target for near targets and insufficiently adapted for far targets (Figure 2D). Diedrichsen and colleagues also showed that force-field adaptation occurs at the same rate with or without concomitant use-dependent learning (Diedrichsen et al., 2010). It appears, therefore, that adaptation is "automatic"; it is an obligate, perhaps reward-indifferent (Mazzoni and Krakauer, 2006), cerebellar-based (Martin et al., 1996a, 1996b; Smith and Shadmehr, 2005; Tseng et al., 2007) learning process that will attempt to reduce prediction errors whenever they occur, even if this is in conflict with task goals.

In spite of the fact that most behavior in error-based motor learning paradigms is well described by adaptation, we argue here that there are phenomena in perturbation paradigms that cannot be explained in terms of adaptation alone. Instead, additional learning mechanisms must be present which are model-free in the sense that they are associated with a memory for action independently of an internal model and are likely to be driven directly by task success (i.e., reward). We posit that there at least two distinct forms of model-free learning processes: use-dependent plasticity, which gives rise to movement biases toward a previously repeated action, and operant reinforcement, which leads to savings when model-based adaptation guides behavior toward a previously successful repeated action.

Combining Model-Based and Model-Free Learning

In the theory of reinforcement learning, the general problem to be solved is to use experience to identify a suitable control policy in an unknown or changing environment (Sutton and Barto, 1998). All motor learning can be conceptualized within this framework; even if there is no explicit reward structure, any task implicitly carries some notion of success or failure that can be encapsulated mathematically through a cost (reward) function.

There are two broad categories of solution methods for such a problem. In a model-based approach, an explicit model of the dynamics of the environment is built from experience, and



this is then used to compute the best possible course of action through standard methods of optimal control theory such as dynamic programming. Note that, in general, model-based control can also entail building a model of the reward structure of the task. In the case of motor control, however, we assume that the reward structure is unambiguous: success is achieved by the cursor reaching the target. In model-free control, by contrast, no such model of the task dynamics is built and instead the value of executing a given action in a given state is learned directly from experience based on subsequent success or failure. While a model-based learning strategy requires significantly less experience to obtain proficient control in an environment and offers greater flexibility (particularly in terms of the ability to generalize knowledge to other tasks), model-free approaches have the advantage of computational simplicity and are not susceptible to problems associated with learning inaccurate or imprecise models (Daw et al., 2005; Dayan, 2009). Therefore, each approach can be advantageous in different circumstances. In sequential discrete decision-making tasks, the brain utilizes both model-based and model-free strategies in parallel (Daw et al., 2005, 2011; Fermin et al., 2010; Gläscher et al., 2010). Theoretical treatments have argued that competition between these two mechanisms enables the benefits of each to be combined to maximum effect (Daw et al., 2005).

Our results suggest that a similar scenario of model-based and model-free learning processes acting in parallel also occurs in the context of motor learning. Adaptation is the model-based component, while model-free components include use-dependent plasticity and operant reinforcement. It is important to note that although the terminology of model-free and model-based learning arises from the theory of reinforcement learning, this does not imply that adaptation is directly sensitive to reward. On the contrary, we believe that adaptation is indifferent to reward outcomes on individual trials, and is purely sensitive to errors in the predicted state of the hand or cursor.

Unlike what has been suggested in the case of sequential decision-making tasks, we believe that under normal circumstances model-based and model-free learning are more cooperative than competitive. In continuous and high-dimensional action spaces, pure model-free learning is unfeasible, especially if a detailed feedback control policy must be acquired. We speculate that during initial learning of a visuomotor rotation, adaptation guides exploration of potential actions toward a suitable solution in hand space, at which point model-free learning becomes more prominent: the asymptotic solution induces use-dependent plasticity through repetition and is reinforced through its operant association with successful adaptation to a perturbation.

Success in a reaching task may not be all-or-nothing, i.e., hitting or missing the target. In fact, we argue that adaptation to errors without actually hitting the target is itself rewarding because it is indicative of imminent success. This idea of the value of "near misses" has been argued for in reinforcement algorithms that assign value to near misses even when actual reinforcement is not given on such trials (MacLin et al., 2007). The rewarding/motivating nature of "near misses" has been reported for gambling where they increase the desire to play (Clark

et al., 2009; Daw et al., 2006; Kakade and Dayan, 2002). Thus, we would argue that movements driven by adaptation are reinforced in hand space because the process of incremental error reduction is the process of ever-closer near misses. Neither repetition alone nor adaptation alone led to savings, which suggests that it is the association of the two that is critical. The novel idea we wish to put forth here is that the association of successful adaptation with a particular movement creates an attractor centered on the movement in hand space. Reexperiencing the same task with the same or even opposite rotational perturbations induces the learner to initially reduce error through pure adaptation but when their movements come within range of the attractor, savings occurs. Furthermore, we conjecture that errors need not be consciously experienced during adaptation in order for the association between the repeated movement and success to occur; all that is required is that adaptation be in operation. There is a precedent for such unconscious reward-based learning in the perceptual learning literature, and the reward can be internal: it does not need to be explicitly provided by the experimenter (Seitz et al., 2009).

Multiple Timescales of Learning

A recent motor learning model has been conceptualized in terms of the existence of fast and slow error-based processes (Kording et al., 2007; Smith et al., 2006). We would argue that skill learning is better conceptualized as cooperation between two qualitatively different kinds of learning: fast model-based adaptation followed by slower improvement through model-free reinforcement. Our previous study of active learning (Huang et al., 2008) in which subjects were allowed to select their own practice sequence to eight targets, each associated with errors of different sizes, can serve as an example of this reconceptualization. We found that subjects repeated successful movements more frequently than error-based learning would predict; from a pure error-based learning perspective, such behavior is suboptimal as it competes with time that could be spent on practice to target directions still associated with large errors - why revisit targets that you have already solved? This behavior is less surprising in our framework, which provides a possible explanation for this apparently sub-optimal behavior; namely that repeating a successful movement is a way to reinforce it. Indeed there are data from other areas of cognitive neuroscience that demonstrate that repeating something that you have successfully learned is the best way to remember it (Chiviacowsky and Wulf, 2007; Karpicke and Roediger, 2008; Wulf and Shea, 2002). We propose that motor skills are acquired through the combination of fast adaptive processes and slower reinforcement processes.

Conclusions

We have shown that use-dependent plasticity and operant reinforcement both occur along with adaptation. Based on our results, we argue that heretofore unexplained, or perhaps erroneously explained, phenomena in adaptation experiments result from the fact that most such experiments inadvertently lie somewhere between our adaptation-only protocol and our adaptation-plus-repetition protocol, with the result that three distinct forms of learning—adaptation, use-dependent plasticity,



and operant reinforcement-are unintentionally lumped together. Future work will need to further dissect these processes and formally model them. The existence of separate learning processes may indicate an underlying anatomical separation. Error-based learning is likely to be cerebellar dependent (Martin et al., 1996a, 1996b; Smith and Shadmehr, 2005; Tseng et al., 2007). Use-dependent learning may occur through Hebbian changes in motor cortex (Orban de Xivry et al., 2011; Verstynen and Sabes, 2011). The presence of dopamine receptors on cells in primary motor cortex (Huntley et al., 1992; Luft and Schwarz, 2009; Ziemann et al., 1997) could provide a candidate mechanism for reward-based modulation of such use-dependent plasticity (Hosp et al., 2011). Our suggestion of an interplay between a model-based process in the cerebellum and a model-free retention process in primary motor cortex is supported by the results of a recent non-invasive brain stimulation study of rotation adaptation; adaptation was accelerated by stimulation of the cerebellum, while stimulation of primary motor cortex led to longer retention (Galea et al., 2010). Finally, operant reinforcement may require dopaminergic projections to the striatum (Wächter et al., 2010). If we are right in our assertion that motor learning studied with error-based paradigms results from the combination of model-free and model-based learning processes then these paradigms may be well suited to study how the brain modularly assembles complex motor abilities.

EXPERIMENTAL PROCEDURES

The Setup

Subjects were seated with their hand and forearm firmly strapped in a splint using padded Velcro bands. The splint was attached to a light-weight frame over a horizontal glass surface. A system of air jets lifted the frame supporting the arm 1 mm above the glass surface, eliminating friction during hand movements. Subjects rested their forehead above the work surface, with their hand and arm hidden from view by a mirror. Targets (green circles) and hand position (indicated, when specified by the task, by a small round cursor) were projected onto the plane of the hand and forearm using a mirror. The arrangement of the mirror, halfway between the hand's workspace and the image formed by the projector, made the virtual images of cursor and targets appear in the same plane as the hand. The workspace was calibrated so that the image of the cursor indicating hand position fell exactly on the unseen tip of the middle finger's location (i.e., veridical display) (Mazzoni et al., 2007).

Hand position was recorded using a pair of 6 degree of freedom magnetic sensors (Flock of Birds, Ascension Technologies, Burlington, VT) placed on the arm and forearm, which transmitted hand position and arm configuration data to the computer at 120 Hz. Custom software recorded hand and arm position in real time and displayed hand position as a cursor on the computer screen. The same software also controlled the display of visual targets.

Subjects

A total of 60 healthy, right-handed subjects participated in the study (mean age = 24.7 ± 4.9 , 25 males). All subjects were naive to the purpose of the study and gave informed consent in compliance to guidelines set forth by the Columbia University Medical Center Institutional Review Board. They were randomly assigned to groups in each experiment.

The Arm Shooting Task

Subjects were asked to make fast, straight, and planar movements through a small circular target displayed veridically using a mirror and monitor (Huang and Shadmehr, 2009; Huang et al., 2008). At the start of a trial, subjects were asked to move the cursor to a starting circle (2.5 mm radius) situated directly in front of them. Once the cursor was in the starting circle, a green, circular target (2.5 mm radius) appeared 6 cm away from the starting circle and the computer

played a short, random-pitch tone, prompting subjects to move. If applicable for the trial, a rotation centered at the starting circle was imposed on the cursor feedback. As soon as the cursor was 6 cm away from the starting circle, a small white dot appeared at the cursor position at that time and remained there for the rest of the trial. Thus, the position of the white dot indicated the angular error the subject made in that trial. Subjects were then asked to return the overshot cursor to the target. The cursor disappeared briefly at this point. Subjects were given feedback regarding movement speed and target accuracy in order to keep these movement variables uniform across individuals. In addition, subjects were verbally encouraged to move faster at the end of a trial if the peak movement speed was less than 80 cm/s. The cursor then reappeared, and subjects brought it back to the starting circle ready for the next trial. All subjects were asked to complete a questionnaire asking them to identify any explicit strategies they might have used during the session.

Experiment 1

Adp⁺Rep⁻ Group

 Adp^+Rep^- subjects (n = 8) performed the reaching task in four types of trial: baseline, training, probe, and washout (Figure 1A). In baseline trials, subjects made movements without additional manipulations to their visual feedback. Targets were randomly chosen from a uniform distribution of directions ranging from 70° to 110° (measured from the positive x axis) totaling 40 possible locations. In training trials, the cursor was rotated counterclockwise (CCW or "+") by a magnitude randomly drawn from a uniform distribution ranging +0° to +40° (Figure S1B). Ten probe trials were interspersed between the 81st and the 160th training trials. These probes were to ten novel targets evenly distributed between 30° to 70° from the positive x axis (Figure 1A). In probe trials, the cursor vanished as soon as it left the starting circle. The washout trials were identical to baseline trials.

Subjects performed these trials in four consecutive blocks with short (1–2 min) breaks between blocks. Block 1 consisted of 80 baseline trials and Block 2, 80 training trials. Block 3 started with 10 probe trials interspersed within 80 training trials and ended with 10 washout trials. Block 4 had 70 washout trials.

Adp⁺Rep⁺ Group

The Adp^+Rep^+ protocol (n = 8) was identical to Adp^+Rep^- except for the order of the imposed rotations in the training trials (Figure 1A). In Adp^+Rep^+ training trials, cursor movements were also rotated by a magnitude drawn from the same distribution as of Adp^+Rep^- training trials (Figure S1B). In Adp^+Rep^+ , however, the optimal movement to cancel out the rotation was always toward the 70° direction (i.e., the repeated direction) in hand space (Figure 1A). For example, the cursor was rotated by +40° when the 110° target was displayed, the rotation was +20° for the 90° target, and +5° for the 75° target, etc. (Figure 1B).

Experiment 2

Adp⁺Rep⁻ and Adp⁺Rep⁺ Groups

 Adp^+Rep^- (n = 10) and Adp^+Rep^+ (n = 10) participated in Experiment 2. The initial training and washout blocks for Adp^+Rep^- and Adp^+Rep^+ in Experiment 2 were identical to their counterparts in Experiment 1 except that training was done without probe trials, and after the washout block, subjects underwent an additional test (relearning) block where they were exposed to a +25° rotation at the 95° target for another 80 trials (Figure 3).

Adp⁻Rep⁻ and Adp⁻Rep⁺ Groups

 Adp^-Rep^- (n = 6) and Adp^-Rep^+ (n = 6) performed the shooting task in three consecutive blocks. In the each of the 160 training trials spanning Block 1 and 2, Adp^-Rep^- aimed for a random target between 70° to 110° without any cursor rotation. In contrast, Adp^-Rep^+ was given only the 70° target in all 160 training trials, also without cursor rotation (Figure 3). Block 3 started with 80 test trials in which both groups were given only the 95° target and their cursor movements were rotated by $+25^\circ$. Forty washout trials immediately followed training with the target relocated to the 70° position and movements were made without cursor rotation.

Experiment 3

$\textbf{SAME-SOLN}_{hand} \ \textbf{and} \ \textbf{SAME-SOLN}_{visual} \ \textbf{Groups}$

SAME-SOLN_{hand} (n = 6) and SAME-SOLN_{visual} (n = 6) groups performed the task in four types of trial: baseline, training, washout, and test trials (Figure 5A).



These two groups performed the task in five consecutive blocks. Block 1 consisted of 80 baseline trials. Block 2 started with 5 baseline trials then followed with 80 training trials. Block 3 began with 80 training trials and finish with 5 baseline trials. Block 4 was a washout block and had 80 baseline trials. Block 4 consisted of 80 test trials (Figure 5A). Baseline and washout trials were the same for both groups and consisted of targets uniformly dispersed between 40° to 100° with no rotation. In training trials, a $+30^{\circ}$ rotation was imposed on a single target. In test trials a -30° rotation was imposed on a single target

In SAME-SOLN_{hand}, the solution in hand space was the same for both training and test trials – arbitrarily chosen to be the movement to the 70° direction in hand space (Figure 5B). Thus, subjects first trained in one target direction (the 100° target) with a +30° rotation and then, after a washout block, trained in another target direction (the 40° target) with a counterrotation of -30° .

In SAME-SOLN_{visual}, the solution in visual/cursor space was the same for both training and test trials (40°) while solutions in hand space were different (Figure 5C). Thus, subjects first trained in one target direction (the 40° target) with a +30° rotation and then, after a washout block, trained to the same target with a -30° rotation.

Data Analysis

Data analysis was performed using Matlab (version R2007a, The Mathworks, Natick, MA). Statistical analysis was performed using SPSS 11.5 (SPSS, Chicago, IL). Unless otherwise specified, t- and p- values were reported using independent-sample 2-tailed t tests. Angular error was calculated as the angular difference between the displayed target center and the white feedback dot. The error reduction rate (i.e., learning and relearning rate) was defined as the time constant obtained by fitting the error time series with a single decaying exponential function of the form $y = C_1 \exp(-rate * x) + C_0$, where C_1 and C_0 are constants, y is the error and x the trial number.

We simulated trial-to-trial hand movement directions in response to the visuomotor rotations as a result of adaptation alone using a single-state state-space model (Donchin et al., 2003; Tanaka et al., 2009). The model equations took the following form:

$$y^{(n)} = R^{(n)} \, - \, K\!\left(T^{(n)}\right) \, z^{(n)}$$

$$z^{(n+1)} = A z^{(n)} + B y^{(n)}$$
.

The k by 1 vector $\mathbf{z}^{(n)}$, is the state of the learner that represents the estimated visuomotor mapping (rotation) associated with each of the k targets in trial n. $K(T^{(n)})$ is the selector matrix that selects the corresponding element in $\mathbf{z}^{(n)}$ for the target $T^{(n)}$. At each trial, $K(T^{(n)})$ $\mathbf{z}^{(n)}$ represents the hand movement direction. The variable R⁽ⁿ⁾ represents the rotation that was imposed; thus, y⁽ⁿ⁾, computed as the difference between R and z, represents the error in the visuomotor mapping (i.e., cursor error). The visuomotor mapping / states of the learner are updated by a generalization function B of size k by 1 that determines how much errors in one target direction affects mapping estimations in neighboring directions. In addition, the visuomotor mapping / states of the learner slowly forget at a rate determined by the scalar A.

To limit the number of parameters in the simulations, we grouped targets in bins with 5° width. Thus, k=16, including all training and probe targets. According to recently published estimations (Tanaka et al., 2009), we interpolated that B, a function of target-to-target angular difference, decreased its gain linearly from 0.09 to 0 within 9 target bins (i.e., a 45° directional window) and that A had a value of 0.98. The motor performance prediction by adaptation alone was simulated deterministically using these parameter values.

Power Analysis

We computed minimum sample sizes on assumed effect sizes for savings based on previously reported data (Zarahn et al., 2008). For an independent samples t test using a two-tailed alpha of 0.05 and power of 0.8, and assuming an effect size d = 1.9375 (computed based on previously reported group means and standard deviation; time constant = 0.47 for savings and 0.16 for naive, with SD = 0.16), the minimum sample size is six subjects per group.

SUPPLEMENTAL INFORMATION

Supplemental Information includes one figure and can be found with this article online at doi:10.1016/j.neuron.2011.04.012.

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REFERENCES

Abeele, S., and Bock, O. (2001). Sensorimotor adaptation to rotated visual input: different mechanisms for small versus large rotations. Exp. Brain Res. 140, 407-410.

Adams, C.D., and Dickinson, A. (1981). Instrumental Responding Following Reinforcer Devaluation. Q. J. Exp. Psychol. B 33, 109-121.

Ajemian, R., D'Ausilio, A., Moorman, H., and Bizzi, E. (2010). Why professional athletes need a prolonged period of warm-up and other peculiarities of human motor learning. J. Mot. Behav. 42, 381-388.

Anderson, M.C., Bjork, E.L., and Bjork, R.A. (2000). Retrieval-induced forgetting: evidence for a recall-specific mechanism. Psychon. Bull. Rev. 7, 522-530.

Arce, F., Novick, I., Mandelblat-Cerf, Y., and Vaadia, E. (2010). Neuronal correlates of memory formation in motor cortex after adaptation to force field. J. Neurosci. 30, 9189-9198.

Avila, I., Reilly, M.P., Sanabria, F., Posadas-Sánchez, D., Chavez, C.L., Banerjee, N., Killeen, P., and Castañeda, E. (2009). Modeling operant behavior in the Parkinsonian rat. Behav. Brain Res. 198, 298-305.

Bédard, P., and Sanes, J.N. (2011). Basal ganglia-dependent processes in recalling learned visual-motor adaptations. Exp. Brain Res. 209, 385-393.

Bock, O., Schneider, S., and Bloomberg, J. (2001). Conditions for interference versus facilitation during sequential sensorimotor adaptation. Exp. Brain Res. 138. 359-365.

Brashers-Krug, T., Shadmehr, R., and Bizzi, E. (1996). Consolidation in human motor memory. Nature 382, 252-255.

Bütefisch, C.M., Davis, B.C., Wise, S.P., Sawaki, L., Kopylev, L., Classen, J., and Cohen, L.G. (2000). Mechanisms of use-dependent plasticity in the human motor cortex. Proc. Natl. Acad. Sci. USA 97, 3661-3665.

Chiviacowsky, S., and Wulf, G. (2007). Feedback after good trials enhances learning. Res. Q. Exerc. Sport 78, 40-47.

Clark, L., Lawrence, A.J., Astley-Jones, F., and Gray, N. (2009). Gambling near-misses enhance motivation to gamble and recruit win-related brain circuitry. Neuron 61, 481-490.

Classen, J., Liepert, J., Wise, S.P., Hallett, M., and Cohen, L.G. (1998). Rapid plasticity of human cortical movement representation induced by practice. J. Neurophysiol. 79, 1117-1123.

Daw, N.D., Niv, Y., and Dayan, P. (2005). Uncertainty-based competition between prefrontal and dorsolateral striatal systems for behavioral control. Nat. Neurosci. 8, 1704-1711.

Daw, N.D., O'Doherty, J.P., Dayan, P., Seymour, B., and Dolan, R.J. (2006). Cortical substrates for exploratory decisions in humans. Nature 441, 876–879.

Daw, N.D., Gershman, S.J., Seymour, B., Dayan, P., and Dolan, R.J. (2011). Model-based influences on humans' choices and striatal prediction errors. Neuron 69, 1204-1215.

Dayan, P. (2009). Goal-directed control and its antipodes. Neural Netw. 22, 213-219.



Debicki, D.B., and Gribble, P.L. (2004). Inter-joint coupling strategy during adaptation to novel viscous loads in human arm movement. J. Neurophysiol. 92, 754–765.

Diedrichsen, J., White, O., Newman, D., and Lally, N. (2010). Use-dependent and error-based learning of motor behaviors. J. Neurosci. *30*, 5159–5166.

Donchin, O., Francis, J.T., and Shadmehr, R. (2003). Quantifying generalization from trial-by-trial behavior of adaptive systems that learn with basis functions: theory and experiments in human motor control. J. Neurosci. 23, 9032–9045.

Doya, K. (1999). What are the computations of the cerebellum, the basal ganglia and the cerebral cortex? Neural Netw. 12, 961–974.

Ebbinghaus, H. (1913). Memory: A Contribution to Experimental Psychology (New York: Teachers College, Columbia University).

Ethier, V., Zee, D.S., and Shadmehr, R. (2008). Spontaneous recovery of motor memory during saccade adaptation. J. Neurophysiol. 99, 2577–2583.

Fermin, A., Yoshida, T., Ito, M., Yoshimoto, J., and Doya, K. (2010). Evidence for model-based action planning in a sequential finger movement task. J. Mot. Behav. 42, 371–379.

Flanagan, J.R., Vetter, P., Johansson, R.S., and Wolpert, D.M. (2003). Prediction precedes control in motor learning. Curr. Biol. *13*, 146–150.

Frank, M.J., Seeberger, L.C., and O'reilly, R.C. (2004). By carrot or by stick: cognitive reinforcement learning in parkinsonism. Science 306, 1940–1943.

Galea, J.M., Vazquez, A., Pasricha, N., Orban de Xivry, J.J., and Celnik, P. (2010). Dissociating the Roles of the Cerebellum and Motor Cortex during Adaptive Learning: The Motor Cortex Retains What the Cerebellum Learns. Cereb. Cortex. Published online December 7, 2010.

Gandolfo, F., Mussa-Ivaldi, F.A., and Bizzi, E. (1996). Motor learning by field approximation. Proc. Natl. Acad. Sci. USA 93, 3843–3846.

Gläscher, J., Daw, N., Dayan, P., and O'Doherty, J.P. (2010). States versus rewards: dissociable neural prediction error signals underlying model-based and model-free reinforcement learning. Neuron 66, 585–595.

Haruno, M., Wolpert, D.M., and Kawato, M. (2001). Mosaic model for sensorimotor learning and control. Neural Comput. 13, 2201–2220.

Held, R., and Rekosh, J. (1963). Motor-sensory feedback and the geometry of visual space. Science 141, 722–723.

Hosp, J.A., Pekanovic, A., Rioult-Pedotti, M.S., and Luft, A.R. (2011). Dopaminergic projections from midbrain to primary motor cortex mediate motor skill learning. J. Neurosci. *31*, 2481–2487.

Huang, V.S., and Shadmehr, R. (2007). Evolution of motor memory during the seconds after observation of motor error. J. Neurophysiol. 97, 3976–3985.

Huang, V.S., and Shadmehr, R. (2009). Persistence of motor memories reflects statistics of the learning event. J. Neurophysiol. *102*, 931–940.

Huang, V.S., Shadmehr, R., and Diedrichsen, J. (2008). Active learning: learning a motor skill without a coach. J. Neurophysiol. *100*, 879–887.

Huntley, G.W., Morrison, J.H., Prikhozhan, A., and Sealfon, S.C. (1992). Localization of multiple dopamine receptor subtype mRNAs in human and monkey motor cortex and striatum. Brain Res. Mol. Brain Res. 15, 181–188.

Imamizu, H., Uno, Y., and Kawato, M. (1995). Internal representations of the motor apparatus: implications from generalization in visuomotor learning. J. Exp. Psychol. Hum. Percept. Perform. *21*, 1174–1198.

Izawa, J., and Shadmehr, R. (2011). Learning from sensory and reward prediction errors during motor adaptation. PLoS Comput. Biol. 7, e1002012.

Jax, S.A., and Rosenbaum, D.A. (2007). Hand path priming in manual obstacle avoidance: evidence that the dorsal stream does not only control visually guided actions in real time. J. Exp. Psychol. Hum. Percept. Perform. 33, 425–441

Joiner, W.M., and Smith, M.A. (2008). Long-term retention explained by a model of short-term learning in the adaptive control of reaching. J. Neurophysiol. *100*, 2948–2955.

Kaelbling, L.P., Littman, M.L., and Moore, A.W. (1996). Reinforcement learning: A survey. J. Artif. Intell. Res. 4, 237–285.

Kakade, S., and Dayan, P. (2002). Dopamine: generalization and bonuses. Neural Netw. 15, 549–559.

Karpicke, J.D., and Roediger, H.L., 3rd. (2008). The critical importance of retrieval for learning. Science 319, 966–968.

Kojima, Y., Iwamoto, Y., and Yoshida, K. (2004). Memory of learning facilitates saccadic adaptation in the monkey. J. Neurosci. 24, 7531–7539.

Kording, K.P., Tenenbaum, J.B., and Shadmehr, R. (2007). The dynamics of memory as a consequence of optimal adaptation to a changing body. Nat. Neurosci. 10, 779–786.

Korenberg, A.T., and Ghahramani, Z. (2002). A Bayesian view of motor adaptation. Current Psychology of Cognition 21, 537–564.

Krakauer, J.W., Ghilardi, M.F., and Ghez, C. (1999). Independent learning of internal models for kinematic and dynamic control of reaching. Nat. Neurosci. 2, 1026–1031.

Krakauer, J.W., Pine, Z.M., Ghilardi, M.F., and Ghez, C. (2000). Learning of visuomotor transformations for vectorial planning of reaching trajectories. J. Neurosci. 20, 8916–8924.

Krakauer, J.W., Ghez, C., and Ghilardi, M.F. (2005). Adaptation to visuomotor transformations: consolidation, interference, and forgetting. J. Neurosci. 25, 473–478

Krutky, M.A., and Perreault, E.J. (2007). Motor cortical measures of use-dependent plasticity are graded from distal to proximal in the human upper limb. J. Neurophysiol. *98*, 3230–3241.

Lackner, J.R., and Dizio, P. (1994). Rapid adaptation to Coriolis force perturbations of arm trajectory. J. Neurophysiol. 72, 299–313.

Lee, J.Y., and Schweighofer, N. (2009). Dual adaptation supports a parallel architecture of motor memory. J. Neurosci. 29, 10396–10404.

Luft, A.R., and Schwarz, S. (2009). Dopaminergic signals in primary motor cortex. Int. J. Dev. Neurosci. 27, 415–421.

MacLeod, M.D., and Macrae, C.N. (2001). Gone but not forgotten: the transient nature of retrieval-induced forgetting. Psychol. Sci. 12, 148–152.

MacLin, O.H., Dixon, M.R., Daugherty, D., and Small, S.L. (2007). Using a computer simulation of three slot machines to investigate a gambler's preference among varying densities of near-miss alternatives. Behav. Res. Methods 39, 237–241.

Malfait, N., Gribble, P.L., and Ostry, D.J. (2005). Generalization of motor learning based on multiple field exposures and local adaptation. J. Neurophysiol. 93, 3327–3338.

Marinelli, L., Crupi, D., Di Rocco, A., Bove, M., Eidelberg, D., Abbruzzese, G., and Ghilardi, M.F. (2009). Learning and consolidation of visuo-motor adaptation in Parkinson's disease. Parkinsonism Relat. Disord. 15, 6–11.

Martin, T.A., Keating, J.G., Goodkin, H.P., Bastian, A.J., and Thach, W.T. (1996a). Throwing while looking through prisms. I. Focal olivocerebellar lesions impair adaptation. Brain *119*, 1183–1198.

Martin, T.A., Keating, J.G., Goodkin, H.P., Bastian, A.J., and Thach, W.T. (1996b). Throwing while looking through prisms. II. Specificity and storage of multiple gaze-throw calibrations. Brain *119*, 1199–1211.

Mazzoni, P., and Krakauer, J.W. (2006). An implicit plan overrides an explicit strategy during visuomotor adaptation. J. Neurosci. 26, 3642–3645.

Mazzoni, P., Hristova, A., and Krakauer, J.W. (2007). Why don't we move faster? Parkinson's disease, movement vigor, and implicit motivation. J. Neurosci. 27, 7105–7116.

Miall, R.C., Jenkinson, N., and Kulkarni, K. (2004). Adaptation to rotated visual feedback: a re-examination of motor interference. Exp. Brain Res. *154*, 201–210.

Orban de Xivry, J.J., Criscimagna-Hemminger, S.E., and Shadmehr, R. (2011). Contributions of the Motor Cortex to Adaptive Control of Reaching Depend on the Perturbation Schedule. Cereb. Cortex. Published online December 3, 2010. 10.1093/cercor/bhq192.

Pine, Z.M., Krakauer, J.W., Gordon, J., and Ghez, C. (1996). Learning of scaling factors and reference axes for reaching movements. Neuroreport 7, 2357–2361.



Rushworth, M.F., Nixon, P.D., Wade, D.T., Renowden, S., and Passingham, R.E. (1998). The left hemisphere and the selection of learned actions. Neuropsychologia 36, 11-24.

Rutledge, R.B., Lazzaro, S.C., Lau, B., Myers, C.E., Gluck, M.A., and Glimcher,P.W. (2009). Dopaminergic drugs modulate learning rates and perseveration in Parkinson's patients in a dynamic foraging task. J. Neurosci. 29, 15104-15114.

Scheidt, R.A., Dingwell, J.B., and Mussa-Ivaldi, F.A. (2001). Learning to move amid uncertainty. J. Neurophysiol. 86, 971-985.

Seitz, A.R., Kim, D., and Watanabe, T. (2009). Rewards evoke learning of unconsciously processed visual stimuli in adult humans. Neuron 61, 700-707. Shadmehr, R., and Mussa-Ivaldi, F.A. (1994). Adaptive representation of dynamics during learning of a motor task. J. Neurosci. 14, 3208-3224.

Shadmehr, R., Smith, M.A., and Krakauer, J.W. (2010). Error correction, sensory prediction, and adaptation in motor control. Annu. Rev. Neurosci. 33, 89-108.

Shohamy, D., Myers, C.E., Grossman, S., Sage, J., and Gluck, M.A. (2005). The role of dopamine in cognitive sequence learning: evidence from Parkinson's disease. Behav. Brain Res. 156, 191-199.

Sing, G.C., and Smith, M.A. (2010). Reduction in learning rates associated with anterograde interference results from interactions between different timescales in motor adaptation. PLoS Comput. Biol. 6, pii: e1000893.

Smith, M.A., and Shadmehr, R. (2005). Intact ability to learn internal models of arm dynamics in Huntington's disease but not cerebellar degeneration. J. Neurophysiol. 93, 2809-2821.

Smith, M.A., Ghazizadeh, A., and Shadmehr, R. (2006). Interacting adaptive processes with different timescales underlie short-term motor learning. PLoS Biol. 4, e179.

Sutton, R.S., and Barto, A.G. (1998). Reinforcement learning: An introduction. In Adaptive Computation and Machine Learning (Cambridge, MA: MIT Press). Tanaka, H., Sejnowski, T.J., and Krakauer, J.W. (2009). Adaptation to visuomotor rotation through interaction between posterior parietal and motor cortical areas. J. Neurophysiol. 102, 2921-2932.

Thoroughman, K.A., and Shadmehr, R. (2000). Learning of action through adaptive combination of motor primitives. Nature 407, 742-747.

Tong, C., Wolpert, D.M., and Flanagan, J.R. (2002). Kinematics and dynamics are not represented independently in motor working memory: evidence from an interference study. J. Neurosci. 22, 1108-1113.

Tseng, Y.W., Diedrichsen, J., Krakauer, J.W., Shadmehr, R., and Bastian, A.J. (2007). Sensory prediction errors drive cerebellum-dependent adaptation of reaching, J. Neurophysiol, 98, 54-62.

Wächter, T., Röhrich, S., Frank, A., Molina-Luna, K., Pekanovic, A., Hertler, B., Schubring-Giese, M., and Luft, A.R. (2010). Motor skill learning depends on protein synthesis in the dorsal striatum after training. Exp. Brain Res. 200,

Wigmore, V., Tong, C., and Flanagan, J.R. (2002). Visuomotor rotations of varying size and direction compete for a single internal model in motor working memory. J. Exp. Psychol. Hum. Percept. Perform. 28, 447-457.

Wixted, J.T. (2004). The psychology and neuroscience of forgetting. Annu. Rev. Psychol. 55, 235-269.

Wulf, G., and Shea, C.H. (2002). Principles derived from the study of simple skills do not generalize to complex skill learning. Psychon. Bull. Rev. 9, 185-211.

Wulf, G., Shea, C., and Lewthwaite, R. (2010). Motor skill learning and performance: a review of influential factors. Med. Educ. 44, 75-84.

Yarrow, K., Brown, P., and Krakauer, J.W. (2009). Inside the brain of an elite athlete: the neural processes that support high achievement in sports. Nat. Rev. Neurosci. 10, 585-596.

Zarahn, E., Weston, G.D., Liang, J., Mazzoni, P., and Krakauer, J.W. (2008). Explaining savings for visuomotor adaptation: linear time-invariant statespace models are not sufficient. J. Neurophysiol. 100, 2537-2548.

Ziemann, U., Tergau, F., Bruns, D., Baudewig, J., and Paulus, W. (1997). Changes in human motor cortex excitability induced by dopaminergic and anti-dopaminergic drugs. Electroencephalogr. Clin. Neurophysiol. 105,

Ziemann, U., Muellbacher, W., Hallett, M., and Cohen, L.G. (2001). Modulation of practice-dependent plasticity in human motor cortex. Brain 124, 1171-

Note Added in Proof

The following manuscript was recently accepted and has been cited several times in our article:

Verstynen, T.D., and Sabes, P.N. (2011). How each movement changes the next: An experimental and theoretical study of fast adaptive priors in reaching. J. Neurosci., in press.